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Patentanmeldung Nr. Patent application No. Demande de brevet n°

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R C van Dijk



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Vironovative B.V.  
Burgemeester Oudlaan 50  
3062 PA Rotterdam  
PAYS-BAS

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Novel atypical pneumonia-causing virus

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**Title: Novel atypical pneumonia-causing virus**

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The invention relates to the field of virology.

The SARS outbreak of 2002-2003 has prompted a search for related viruses that may have previously caused atypical pneumonias or that may do so in the future. A 10 respiratory illness (atypical pneumonia) was diagnosed in an 8 months old patient that could not be attributed to SARS (Severe Acute Respiratory Syndrome) virus or any other known viral infection. The patient tested negative for influenza, parainfluenza, mumps and RSV and yet the disease was identified to be caused by a virus which closely resembled SARS.

15 For being able to trace its origin, monitor its epidemiology and prevent possible spreading of the disease, it is of great importance to be able to recognise viral causes of pneumonia in an early stage. Especially, if severe diseases are found to be caused by viruses, it is necessary to detect the identity of the virus as soon as possible, in order to develop diagnostic tools and possibly therapies. The SARS epidemic has shown that it is 20 paramount for prevention of spread of the disease to be able to get an early diagnosis in order to take timely and effective isolation measures and initiate quarantine precautions. Only then, world-wide contaminations can be prevented.

Furthermore, identification of the viral cause for the disease enables development of vaccines, which can be used prophylactically to protect people who are a 25 risk of being infected. And, finally, knowledge of the viral cause enables to develop therapeutic measures.

Thus, there is great need in developing diagnostic tools and therapies for viral pneumonias in general, and particular to a novel disease-causing infectious agent, especially when this agent appears to be a virus.

30 The invention provides the nucleotide sequence of an isolated essentially mammalian positive-sense single stranded RNA virus belonging to the Coronaviruses, which is the causative factor for the new disease, hereinafter referred to as EMCR-CoV and the disease being referred to as EMCR-CoV-caused pneumonia. A virus according to

the invention is isolatable from a human with respiratory tract disease such as, but not limited to, atypical pneumonia.

From a phylogenetic analysis of the Matrix and Nucleocapsid gene sequences of the virus (Fig. 1a and 1b) it appears that the virus is a distinct member of the group 5 formed by PEDV (porcine epidemic diarrhea virus), HCoV-229E (human coronavirus 229E), PRCoV (porcine respiratory coronavirus), TGEV (transmissible gastroenteritis virus), CaCoV (Canine coronavirus) and FeCoV (feline coronavirus). In general, human coronavirus 229E seems to be the closest relative (at least for the Matrix and Nucleocapsid proteins).

10 Although phylogenetic analyses provide a convenient method of identifying a virus, several other possibly more straightforward albeit somewhat more coarse methods for identifying said virus or viral proteins or nucleic acids from said virus are herein also provided. As a rule of thumb an EMCR-Coronavirus can be identified by the percentages of homology of the virus, proteins or nucleic acids to be identified in 15 comparison with viral proteins or nucleic acids identified herein by sequence. It is generally known that virus species, especially RNA virus species, often constitute a quasi species wherein a cluster of said viruses displays heterogeneity among its members. Thus it is expected that each isolate may have a somewhat different percentage relationship with the sequences of the isolate as provided herein.

20 When one wishes to compare a virus isolate with the sequences as listed in figure 3, the invention provides an isolated essentially mammalian positive-sense single stranded RNA virus (EMCR-CoV) belonging to the Coronaviruses and identifiable as phylogenetically corresponding thereto by determining a nucleic acid sequence of said virus and determining that said nucleic acid sequence has a percentage nucleic acid 25 identity to the sequences as listed higher than the percentages identified herein for the nucleic acids as identified herein below in comparison with PEDV, 229E, PRCoV, TGEV CaCoV and FeCoV. Likewise, an isolated essentially mammalian positive-sense single stranded RNA virus (EMCR-CoV) belonging to the Coronaviruses and identifiable as phylogenetically corresponding thereto by determining an amino acid sequence of said 30 virus and determining that said amino acid sequence has a percentage amino acid homology to the sequences as listed which is essentially higher than the percentages provided herein in comparison with PEDV, 229E, PRCoV, TGEV, CaCoV and FeCoV.

With the provision of the sequence information of this EMCR-Coronavirus (EMCR-CoV), the invention provides diagnostic means and methods, prophylactic mean

and methods and therapeutic means and methods to be employed in the diagnosis, prevention and/or treatment of disease, in particular of respiratory disease (atypical pneumonia), in particular of mammals, more in particular in humans associated with infection by this virus. In virology, it is most advisory that diagnosis, prophylaxis and/or treatment of a specific viral infection is performed with reagents that are most specific for said specific virus causing said infection. In this case this means that it is preferred that said diagnosis, prophylaxis and/or treatment of an EMC-CoV virus infection is performed with reagents that are most specific for EMC-CoV virus. This by no means however excludes the possibility that less specific, but sufficiently cross-reactive reagents are used instead, for example because they are more easily available and sufficiently address the task at hand.

The invention for example provides a method for virologically diagnosing an EMC-CoV infection of an animal, in particular of a mammal, more in particular of a human being, comprising determining in a sample of said animal the presence of a viral isolate or component thereof by reacting said sample with an EMC-CoV specific nucleic acid or antibody according to the invention, and a method for serologically diagnosing an EMC-CoV infection of a mammal comprising determining in a sample of said mammal the presence of an antibody specifically directed against an EMC-CoV virus or component thereof by reacting said sample with an EMC-CoV virus-specific proteinaceous molecule or fragment thereof or an antigen according to the invention.

The invention also provides a diagnostic kit for diagnosing an EMC-CoV infection comprising an EMC-CoV virus, an EMC-CoV virus-specific nucleic acid, proteinaceous molecule or fragment thereof, antigen and/or an antibody according to the invention, and preferably a means for detecting said EMC-CoV virus, EMC-CoV virus-specific nucleic acid, proteinaceous molecule or fragment thereof, antigen and/or an antibody, said means for example comprising an excitable group such as a fluorophore or enzymatic detection system used in the art (examples of suitable diagnostic kit format comprise IF, ELISA, neutralization assay, RT-PCR assay). To determine whether an as yet unidentified virus component or synthetic analogue thereof such as nucleic acid, proteinaceous molecule or fragment thereof can be identified as EMC-CoV-virus-specific, it suffices to analyse the nucleic acid or amino acid sequence of said component, for example for a stretch of said nucleic acid or amino acid, preferably of at least 10, more preferably at least 25, more preferably at least 40 nucleotides or amino acids (respectively), by sequence homology comparison with the

provided EMCR-CoV viral sequences and with known non-EMCR-CoV viral sequences (human coronavirus 299E is preferably used) using for example phylogenetic analyses as provided herein. Depending on the degree of relationship with said EMCR-CoV or non-EMCR-CoV viral sequences, the component or synthetic analogue can be identified.

5 The invention thus provides the nucleotide sequence of a novel etiological agent, an isolated essentially mammalian positive-sense single stranded RNA virus (herein also called EMCR-CoV virus) belonging to the Coronaviridae family, and EMCR-CoV virus-specific components or synthetic analogues thereof.

Coronaviruses were first isolated from chickens in 1937, while the first human 10 coronavirus was propagated *in vitro* by Tyrell and Bonoe in 1965. There are now about 13 species in this family, which infect cattle, pigs, rodents, cats, dogs, birds and man. Coronavirus particles are irregularly shaped, about 60-220 nm in diameter, with an outer envelope bearing distinctive, 'club-shaped' peplomers ( about 20 nm long and 10 nm wide at the distal end). This 'crown-like' appearance give the family its name. The 15 envelope carries two glycoproteins: S, the spike glycoprotein which is involved in cell fusion and is a major antigen, and M, the membrane glycoprotein, which is involved in budding and envelope formation. The genome is associated with a basic phosphoprotein, designated N. The genome of coronaviruses, a single stranded positive-sense RNA strand, is typically 27-31 Kb long and contains a 5' methylated cap and a 3' poly-A tail, 20 by which it can directly function as an mRNA in the infected cell. Initially the 5' ORF 1 (about 20 Kb) is translated to produce a viral polymerase, which then produces a full length negative sense strand. This is used as a template to produce mRNA as a 'nested set' of transcripts, all with identical 5' non-translated leader sequence of 72 nucleotides and coincident 3' polyadenylated ends. Each mRNA thus produced is monocistronic, the 25 genes at the 5' end being translated from the longest mRNA and so on. These unusual cytoplasmic structures are produced not by splicing, but by the polymerase during transcription. Between each of the genes there is a repeated intergenic sequence – AACUAAAC – which interacts with the transcriptase plus cellular factors to splice the leader sequence onto the start of each ORF. In some coronaviruses there are about 8 30 ORFs, coding for the proteins mentioned above, but also for a haemagglutinin esterase (HE), and several other non-structural proteins.

Newly isolated viruses are phylogenetically corresponding to and thus taxonomically corresponding to EMCR-CoV virus when comprising a gene order and/or amino acid sequence and/or nucleotide sequence sufficiently similar to our prototypic

EMCR-CoV virus. The highest amino acid sequence homology, between EMCR-CoV virus and any of the known other viruses of the same family to date (human coronavirus 299E or Porcine Epidemic Diarrhea Virus) is for parts of the replicase polyprotein 1ab 80-88% (see, for example Fig. 3 sequences D and E; the % homology, and the virus to which the homology is found depend on the region of the replicase that is examined), as can be deduced when comparing the sequences given in figure 3 with sequences of other viruses, in particular of human coronavirus 299E. Individual proteins or whole virus isolates with, respectively, higher homology than these mentioned maximum values are considered phylogenetically corresponding and thus taxonomically corresponding to EMCR-CoV virus, and generally will be encoded by a nucleic acid sequence structurally corresponding with a sequence as shown in figure 3. Herewith the invention provides a virus phylogenetically corresponding to the isolated virus of which the sequences are depicted in figure 3.

It should be noted that, similar to other viruses, a certain degree of variation can be expected to be found between EMCR-CoV-viruses isolated from different sources.

Also, the viral sequence of the EMCR-CoV virus or an isolated EMCR-CoV virus gene as provided herein for example shows less than 95%, preferably less than 90%, more preferably less than 80%, more preferably less than 70% and most preferably less than 65% nucleotide sequence homology or less than 95%, preferably less than 90%, more preferably less than 80%, more preferably less than 70% and most preferably less than 65% amino acid sequence homology with the respective nucleotide or amino acid sequence of the human coronavirus 299E or Porcine Epidemic Diarrhea Virus as for example can be found in Genbank (for example in accession number af304460 (HCoV-299E) or af353511 (PEDV)).

Sequence divergence of EMCR-CoV strains around the world may be somewhat higher, in analogy with other coronaviruses.

The term "nucleotide sequence homology" as used herein denotes the presence of homology between two (poly)nucleotides. Polynucleotides have "homologous" sequences if the sequence of nucleotides in the two sequences is the same when aligned for maximum correspondence. Sequence comparison between two or more polynucleotides is generally performed by comparing portions of the two sequences over a comparison window to identify and compare local regions of sequence similarity. The comparison window is generally from about 20 to 200 contiguous nucleotides. The "percentage of sequence homology" for polynucleotides, such as 50, 60, 70, 80, 90, 95, 98, 99 or 100

percent sequence homology may be determined by comparing two optimally aligned sequences over a comparison window, wherein the portion of the polynucleotide sequence in the comparison window may include additions or deletions (i.e. gaps) as compared to the reference sequence (which does not comprise additions or deletions) for 5 optimal alignment of the two sequences. The percentage is calculated by: (a) determining the number of positions at which the identical nucleic acid base occurs in both sequences to yield the number of matched positions; (b) dividing the number of matched positions by the total number of positions in the window of comparison; and (c) multiplying the result by 100 to yield the percentage of sequence homology. Optimal 10 alignment of sequences for comparison may be conducted by computerized implementations of known algorithms, or by inspection. Readily available sequence comparison and multiple sequence alignment algorithms are, respectively, the Basic Local Alignment Search Tool (BLAST) (Altschul, S.F. et al. 1990. J. Mol. Biol. 215:403; Altschul, S.F. et al. 1997. Nucleic Acid Res. 25:3389-3402) and ClustalW programs both 15 available on the internet. Other suitable programs include GAP, BESTFIT and FASTA in the Wisconsin Genetics Software Package (Genetics Computer Group (GCG), Madison, WI, USA).

As used herein, "substantially complementary" means that two nucleic acid sequences have at least about 65%, preferably about 70%, more preferably about 80%, even more 20 preferably 90%, and most preferably about 98%, sequence complementarity to each other. This means that the primers and probes must exhibit sufficient complementarity to their template and target nucleic acid, respectively, to hybridise under stringent conditions. Therefore, the primer sequences as disclosed in this specification need not reflect the exact sequence of the binding region on the template and degenerate primers 25 can be used. A substantially complementary primer sequence is one that has sufficient sequence complementarity to the amplification template to result in primer binding and second-strand synthesis.

The term "hybrid" refers to a double-stranded nucleic acid molecule, or duplex, formed by hydrogen bonding between complementary nucleotides. The terms "hybridise" 30 or "anneal" refer to the process by which single strands of nucleic acid sequences form double-helical segments through hydrogen bonding between complementary nucleotides.

The term "oligonucleotide" refers to a short sequence of nucleotide monomers (usually 6 to 100 nucleotides) joined by phosphorous linkages (e.g., phosphodiester, alkyl and aryl-phosphate, phosphorothioate), or non-phosphorous linkages (e.g., peptide,

sulfamate and others). An oligonucleotide may contain modified nucleotides having modified bases (e.g., 5-methyl cytosine) and modified sugar groups (e.g., 2'-O-methyl ribosyl, 2'-O-methoxyethyl ribosyl, 2'-fluoro ribosyl, 2'-amino ribosyl, and the like). Oligonucleotides may be naturally-occurring or synthetic molecules of double- and

5 single-stranded DNA and double- and single-stranded RNA with circular, branched or linear shapes and optionally including domains capable of forming stable secondary structures (e.g., stem-and-loop and loop-stem-loop structures).

The term "primer" as used herein refers to an oligonucleotide which is capable of annealing to the amplification target allowing a DNA polymerase to attach thereby

10 serving as a point of initiation of DNA synthesis when placed under conditions in which synthesis of primer extension product which is complementary to a nucleic acid strand is induced, i.e., in the presence of nucleotides and an agent for polymerization such as DNA polymerase and at a suitable temperature and pH. The (amplification) primer is preferably single stranded for maximum efficiency in amplification. Preferably, the

15 primer is an oligodeoxy ribonucleotide. The primer must be sufficiently long to prime the synthesis of extension products in the presence of the agent for polymerization. The exact lengths of the primers will depend on many factors, including temperature and source of primer. A "pair of bi-directional primers" as used herein refers to one forward and one reverse primer as commonly used in the art of DNA amplification such as in

20 PCR amplification.

The term "probe" refers to a single-stranded oligonucleotide sequence that will recognize and form a hydrogen-bonded duplex with a complementary sequence in a target nucleic acid sequence analyte or its cDNA derivative.

The terms "stringency" or "stringent hybridization conditions" refer to

25 hybridization conditions that affect the stability of hybrids, e.g., temperature, salt concentration, pH, formamide concentration and the like. These conditions are empirically optimised to maximize specific binding and minimize non-specific binding of primer or probe to its target nucleic acid sequence. The terms as used include reference to conditions under which a probe or primer will hybridise to its target sequence, to a

30 detectably greater degree than other sequences (e.g. at least 2-fold over background). Stringent conditions are sequence dependent and will be different in different circumstances. Longer sequences hybridise specifically at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (T<sub>m</sub>) for the specific sequence at a defined ionic strength and pH. The T<sub>m</sub>

is the temperature (under defined ionic strength and pH) at which 50% of a complementary target sequence hybridises to a perfectly matched probe or primer. Typically, stringent conditions will be those in which the salt concentration is less than about 1.0 M Na<sup>+</sup> ion, typically about 0.01 to 1.0 M Na<sup>+</sup> ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes or primers (e.g. 10 to 50 nucleotides) and at least about 60°C for long probes or primers (e.g. greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. Exemplary low stringent conditions or "conditions of reduced stringency" include hybridization with a buffer solution of 30% formamide, 1 M NaCl, 1% SDS at 37°C and a wash in 2x SSC at 40°C. Exemplary high stringency conditions include hybridization in 50% formamide, 1 M NaCl, 1% SDS at 37°C, and a wash in 0.1x SSC at 60°C. Hybridization procedures are well known in the art and are described in e.g. Ausubel et al, Current Protocols in Molecular Biology, John Wiley & Sons Inc., 1994.

The term "antibody" includes reference to antigen binding forms of antibodies (e.g., Fab, F (ab) 2). The term "antibody" frequently refers to a polypeptide substantially encoded by an immunoglobulin gene or immunoglobulin genes, or fragments thereof which specifically bind and recognize an analyte (antigen). However, while various antibody fragments can be defined in terms of the digestion of an intact antibody, one of skill will appreciate that such fragments may be synthesized de novo either chemically or by utilizing recombinant DNA methodology. Thus, the term antibody, as used herein, also includes antibody fragments such as single chain Fv, chimeric antibodies (i. e., comprising constant and variable regions from different species), humanized antibodies (i. e., comprising a complementarity determining region (CDR) from a non-human source) and heteroconjugate antibodies (e. g., bispecific antibodies).

In short, the invention provides an isolated essentially mammalian positive-sense single stranded RNA virus (EMCR-CoV) belonging to the Coronaviruses and identifiable as phylogenetically corresponding thereto by determining a nucleic acid sequence of a suitable fragment of the genome of said virus and testing it in phylogenetic tree analyses wherein maximum likelihood trees are generated using 100 bootstraps and 3 jumbles and finding it to be more closely phylogenetically corresponding to a virus isolate having the sequences as depicted in figure 3 than it is corresponding to a virus isolate of PEDV (porcine epidemic diarrhea virus), HCoV-229E (human coronavirus 229E), PRCoV (porcine respiratory coronavirus), TGEV

(transmissible gastroenteritis virus), CaCoV (Canine coronavirus) and FeCoV (feline coronavirus).

Suitable nucleic acid genome fragments each useful for such phylogenetic tree analyses are for example any of the fragments encoding the Matrix protein or the 5 Nucleocapsid protein as disclosed in figure 3, leading to the phylogenetic tree analysis as disclosed herein in figure 1a or 1b.

A suitable open reading frame (ORF) comprises the ORF encoding the viral replicate (ORF 1a). When an overall amino acid identity of at least 60%, preferably of at least 70%, more preferably of at least 80%, more preferably of at least 90%, most 10 preferably of at least 95% of the analysed replicate with the replicate having a sequence comprising the amino acid fragments A, B, C, D, E, and/or F of figure 3 is found, the analysed virus isolate comprises an EMC-CoV virus isolate according to the invention.

Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the Nucleocapsid protein. When an overall amino acid 15 identity of at least 60%, more preferably of at least 70%, more preferably of at least 80%, more preferably of at least 90%, most preferably of at least 95% of the analysed Nucleocapsid protein with the Nucleocapsid protein encoded by a sequence comprising (part of) the sequence F of figure 3 is found, the analysed virus isolate comprises an EMC-CoV isolate according to the invention.

20 Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the Matrix protein. When an overall amino acid identity of at least 60%, more preferably of at least 70%, more preferably of at least 80%, more preferably of at least 90%, most preferably of at least 95% of the analysed Matrix protein with the Matrix protein encoded by a sequence comprising (part of) the sequence 25 F of figure 3 is found, the analysed virus isolate comprises an EMC-CoV isolate according to the invention.

Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the spike protein S. When an overall amino acid identity of 30 at least 60%, more preferably of at least 70%, more preferably of at least 80%, more preferably of at least 90%, most preferably of at least 95% of the analysed S-protein encoded by a sequence comprising the sequence of translation 2 of E and translation 1 of the F sequence of the S-protein as depicted in figure 3 is found, the analysed virus isolate comprises an EMC-CoV virus isolate according to the invention. The S ORF of the EMC-CoV virus seems to be located adjacent to the ORF 1ab (coding for the viral

replicase), which would discriminate an EMCR-CoV viruses from the bovine coronavirus and the murine hepatitis virus, which have a so-called 2a gene and an HE-gene between the S protein and the viral polymerase.

5 The invention provides among others an isolated or recombinant nucleic acid or virus-specific functional fragment thereof obtainable from a virus according to the invention. The isolated or recombinant nucleic acids comprises the sequences as given in figure 3 or sequences of homologues which are able to hybridise with those under stringent conditions. In particular, the invention provides primers and/or probes suitable for identifying an EMCR-CoV virus nucleic acid.

10 Furthermore, the invention provides a vector comprising a nucleic acid according to the invention. To begin with, vectors such as plasmid vectors containing (parts of) the genome of the EMCR-CoV virus, virus vectors containing (parts of) the genome of the EMCR-CoV (for example, but not limited thereto, vaccinia virus, retroviruses, baculovirus), or EMCR-CoV virus containing (parts of) the genome of other viruses or 15 other pathogens are provided.

Also, the invention provides a host cell comprising a nucleic acid or a vector according to the invention. Plasmid or viral vectors containing the replicase components of EMCR-CoV virus are generated in prokaryotic cells for the expression of the components in relevant cell types (bacteria, insect cells, eukaryotic cells). Plasmid or 20 viral vectors containing full-length or partial copies of the EMCR-CoV virus genome will be generated in prokaryotic cells for the expression of viral nucleic acids *in-vitro* or *in-vivo*. The latter vectors may contain other viral sequences for the generation of chimeric viruses or chimeric virus proteins, may lack parts of the viral genome for the generation of replication defective virus, and may contain mutations, deletions or insertions for the 25 generation of attenuated viruses.

Infectious copies of EMCR-CoV virus (being wild type, attenuated, replication-defective or chimeric) can be produced upon co-expression of the polymerase components according to the state-of-the-art technologies described above.

30 In addition, eukaryotic cells, transiently or stably expressing one or more full-length or partial EMCR-CoV virus proteins can be used. Such cells can be made by transfection (proteins or nucleic acid vectors), infection (viral vectors) or transduction (viral vectors) and may be useful for complementation of mentioned wild type, attenuated, replication-defective or chimeric viruses.

A chimeric virus may be of particular use for the generation of recombinant vaccines protecting against two or more viruses. For example, it can be envisaged that EMCR-CoV virus vector expressing one or more proteins of a human metapneumovirus or a human metapneumovirus vector expressing one or more proteins of EMCR-CoV

5 virus will protect individuals vaccinated with such vector against both virus infections. Such a specific chimeric virus is particularly useful in the invention because it is suspected that co-infection of, for instance, human metapneumovirus frequently occurs in coronavirus infected patients. Attenuated and replication-defective viruses may be of use for vaccination purposes with live vaccines as has been suggested for other viruses.

10 In a preferred embodiment, the invention provides a proteinaceous molecule or coronavirus-specific viral protein or functional fragment thereof encoded by a nucleic acid according to the invention. Useful proteinaceous molecules are for example derived from any of the genes or genomic fragments derivable from a virus according to the invention. Such molecules, or antigenic fragments thereof, as provided herein, are for

15 example useful in diagnostic methods or kits and in pharmaceutical compositions such as sub-unit vaccines and inhibitory peptides. Particularly useful are the viral replicase protein, the spike protein, the matrix protein, the nucleocapsid or antigenic fragments thereof for inclusion as antigen or subunit immunogen, but inactivated whole virus can also be used. Particulary useful are also those proteinaceous substances that are

20 encoded by recombinant nucleic acid fragments that are identified for phylogenetic analyses, of course preferred are those that are within the preferred bounds and metes of ORFs useful in phylogenetic analyses, in particular for eliciting EMCR-CoV virus specific antibodies, whether *in vivo* (e.g. for protective puposes or for providing diagnostic antibodies) or *in vitro* (e.g. by phage display technology or another technique

25 useful for generating synthetic antibodies).

Also provided herein are antibodies, be it natural polyclonal or monoclonal, or synthetic (e.g. (phage) library-derived binding molecules) antibodies that specifically react with an antigen comprising a proteinaceous molecule or EMCR-CoV virus-specific functional fragment thereof according to the invention. Such antibodies are useful in a

30 method for identifying a viral isolate as an EMCR-CoV virus comprising reacting said viral isolate or a component thereof with an antibody as provided herein. This can for example be achieved by using purified or non-purified EMCR-CoV virus or parts thereof (proteins, peptides) using ELISA, RIA, FACS or similar formats of antigen detection assays (Current Protocols in Immunology). Alternatively, infected cells or cell cultures

may be used to identify viral antigens using classical immunofluorescence or immunohistochemical techniques. Specifically useful in this respect are antibodies raised against EMCR-CoV virus proteins which are encoded by a nucleotide sequence comprising one or more of the fragments disclosed in figure 3.

5 Other methods for identifying a viral isolate as an EMCR-CoV virus comprise reacting said viral isolate or a component thereof with a virus specific nucleic acid according to the invention.

10 In this way the invention provides a viral isolate identifiable with a method according to the invention as a mammalian virus taxonomically corresponding to a positive-sense single stranded RNA virus identifiable as likely belonging to the EMCR-CoV virus genus within the family of Coronaviruses.

15 The method is useful in a method for virologically diagnosing an EMCR-CoV virus infection of a mammal, said method for example comprising determining in a sample of said mammal the presence of a viral isolate or component thereof by reacting said sample with a nucleic acid or an antibody according to the invention.

Methods of the invention can in principle be performed by using any nucleic acid amplification method, such as the Polymerase Chain Reaction (PCR; Mullis 1987, U.S. Pat. No. 4,683,195, 4,683,202, en 4,800,159) or by using amplification reactions such as Ligase Chain Reaction (LCR; Barany 1991, Proc. Natl. Acad. Sci. USA 88:189-193; EP Appl. No., 320,308), Self-Sustained Sequence Replication (3SR; Guatelli et al., 1990, Proc. Natl. Acad. Sci. USA 87:1874-1878), Strand Displacement Amplification (SDA; U.S. Pat. Nos. 5,270,184, en 5,455,166), Transcriptional Amplification System (TAS; Kwoh et al., Proc. Natl. Acad. Sci. USA 86:1173-1177), Q-Beta Replicase (Lizardi et al., 1988, Bio/Technology 6:1197), Rolling Circle Amplification (RCA; U.S. Pat. No. 5,871,921), Nucleic Acid Sequence Based Amplification (NASBA), Cleavase Fragment Length Polymorphism (U.S. Pat. No. 5,719,028), Isothermal and Chimeric Primer-initiated Amplification of Nucleic Acid (ICAN), Ramification-extension Amplification Method (RAM; U.S. Pat. Nos. 5,719,028 and 5,942,391) or other suitable methods for amplification of nucleic acids.

30 In order to amplify a nucleic acid with a small number of mismatches to one or more of the amplification primers, an amplification reaction may be performed under conditions of reduced stringency (e.g. a PCR amplification using an annealing temperature of 38°C, or the presence of 3.5 mM MgCl<sub>2</sub>). The person skilled in the art will be able to select conditions of suitable stringency.

The primers herein are selected to be "substantially" complementary (i.e. at least 65%, more preferably at least 80% perfectly complementary) to their target regions present on the different strands of each specific sequence to be amplified. It is possible to use primer sequences containing e.g. inositol residues or ambiguous bases or even 5 primers that contain one or more mismatches when compared to the target sequence. In general, sequences that exhibit at least 65%, more preferably at least 80% homology with the target DNA or RNA oligonucleotide sequences, are considered suitable for use in a method of the present invention. Sequence mismatches are also not critical when using low stringency hybridization conditions.

10 The detection of the amplification products can in principle be accomplished by any suitable method known in the art. The detection fragments may be directly stained or labelled with radioactive labels, antibodies, luminescent dyes, fluorescent dyes, or enzyme reagents. Direct DNA stains include for example intercalating dyes such as acridine orange, ethidium bromide, ethidium monoazide or Hoechst dyes.

15 Alternatively, the DNA or RNA fragments may be detected by incorporation of labelled dNTP bases into the synthesized fragments. Detection labels which may be associated with nucleotide bases include e.g. fluorescein, cyanine dye or BrdUrd.

When using a probe-based detection system, a suitable detection procedure for 20 use in the present invention may for example comprise an enzyme immunoassay (EIA) format (Jacobs et al., 1997, J. Clin. Microbiol. 35, 791-795). For performing a detection by manner of the EIA procedure, either the forward or the reverse primer used in the amplification reaction may comprise a capturing group, such as a biotin group for immobilization of target DNA PCR amplicons on e.g. a streptavidin coated microtiter plate wells for subsequent EIA detection of target DNA -amplicons (see below). The 25 skilled person will understand that other groups for immobilization of target DNA PCR amplicons in an EIA format may be employed.

30 Probes useful for the detection of the target DNA as disclosed herein preferably bind only to at least a part of the DNA sequence region as amplified by the DNA amplification procedure. Those of skill in the art can prepare suitable probes for detection based on the nucleotide sequence of the target DNA without undue experimentation as set out herein. Also the complementary nucleotide sequences, whether DNA or RNA or chemically synthesized analogs, of the target DNA may suitably be used as type-specific detection probes in a method of the invention, provided that such a complementary strand is amplified in the amplification reaction employed.

Suitable detection procedures for use herein may for example comprise immobilization of the amplicons and probing the DNA sequences thereof by e.g. southern blotting. Other formats may comprise an EIA format as described above. To facilitate the detection of binding, the specific amplicon detection probes may comprise a 5 label moiety such as a fluorophore, a chromophore, an enzyme or a radio-label, so as to facilitate monitoring of binding of the probes to the reaction product of the amplification reaction. Such labels are well-known to those skilled in the art and include, for example, fluorescein isothiocyanate (FITC),  $\beta$ -galactosidase, horseradish peroxidase, streptavidin, biotin, digoxigenin,  $^{35}\text{S}$  or  $^{125}\text{I}$ . Other examples will be apparent to those skilled in the 10 art.

15 Detection may also be performed by a so called reverse line blot (RLB) assay, such as for instance described by Van den Brule et al. (2002, *J. Clin. Microbiol.* 40, 779-787). For this purpose RLB probes are preferably synthesized with a 5' amino group for subsequent immobilization on e.g. carboxyl-coated nylon membranes. The advantage of an RLB format is the ease of the system and its speed, thus allowing for high throughput sample processing.

20 The use of nucleic acid probes for the detection of RNA or DNA fragments is well known in the art. Mostly these procedure comprise the hybridization of the target nucleic acid with the probe followed by post-hybridization washings. Specificity is typically the function of post-hybridization washes, the critical factors being the ionic strength and temperature of the final wash solution. For nucleic acid hybrids, the  $\text{Tm}$  can be approximated from the equation of Meinkoth and Wahl, *Anal. Biochem.*, 138: 267-284 (1984):  $\text{Tm} = 81.5 \text{ }^{\circ}\text{C} + 16.6 (\log M) + 0.41 (\% \text{ GC}) - 0.61 (\% \text{ form}) - 500/L$ ; where  $M$  is the molarity of monovalent cations, % GC is the percentage of guanosine and cytosine 25 nucleotides in the nucleic acid, % form is the percentage of formamide in the hybridization solution, and  $L$  is the length of the hybrid in base pairs. The  $\text{Tm}$  is the temperature (under defined ionic strength and pH) at which 50% of a complementary target sequence hybridizes to a perfectly matched probe.  $\text{Tm}$  is reduced by about 1  $^{\circ}\text{C}$  for each 1 % of mismatching; thus, the hybridization and/or wash conditions can be 30 adjusted to hybridize to sequences of the desired identity. For example, if sequences with > 90% identity are sought, the  $\text{Tm}$  can be decreased 10  $^{\circ}\text{C}$ . Generally, stringent conditions are selected to be about 5  $^{\circ}\text{C}$  lower than the thermal melting point ( $\text{Tm}$ ) for the specific sequence and its complement at a defined ionic strength and pH. However, severely stringent conditions can utilize a hybridization and/or wash at 1,2,3, or 4  $^{\circ}\text{C}$

lower than the thermal melting point (Tm); moderately stringent conditions can utilize hybridization and/or wash at 6, 7, 8, 9, or 10 °C lower than the thermal melting point (Tm); low stringency conditions can utilize a hybridization and/or wash at 11, 12, 13, 14, 15, or 20 °C lower than the thermal melting point (Tm). Using the equation,

5 hybridization and wash compositions, and desired Tm, those of ordinary skill will understand that variations in the stringency of hybridization and/or wash solutions are inherently described. If the desired degree of mismatching results in a Tm of less than 45 °C (aqueous solution) or 32 °C (formamide solution) it is preferred to increase the SSC concentration so that a higher temperature can be used. An extensive guide to the

10 hybridization of nucleic acids is found in Tijssen, *Laboratory Techniques in Biochemistry and Molecular Biology—Hybridization with Nucleic Acid Probes, Part I, Chapter 2*" Overview of principles of hybridization and the strategy of nucleic acid probe assays", Elsevier, New York (1993); and *Current Protocols in Molecular Biology, Chapter 2*, Ausubel, et al., Eds., Greene Publishing and Wiley-Interscience, New York (1995).

15 In another aspect, the invention provides oligonucleotide probes for the generic detection of target RNA or DNA. The detection probes herein are selected to be "substantially" complementary to one of the strands of the double stranded nucleic acid generated by an amplification reaction of the invention. Preferably the probes are substantially complementary to the immobilizable, e.g. biotin labelled, antisense strand

20 of the amplicons generated from the target RNA or DNA.

It is allowable for detection probes of the present invention to contain one or more mismatches to their target sequence. In general, sequences that exhibit at least 65%, more preferably at least 80% homology with the target oligonucleotide sequences are considered suitable for use in a method of the present invention.

25 Antibodies, both monoclonal and polyclonal, can also be used for detection purpose in the present invention, for example, in immunoassays in which they can be utilized in liquid phase or bound to a solid phase carrier. In addition, the monoclonal antibodies in these immunoassays can be detectably labeled in various ways. A variety of immunoassay formats may be used to select antibodies specifically reactive with a

30 particular protein (or other analyte). For example, solid-phase ELISA immunoassays are routinely used to select monoclonal antibodies specifically immunoreactive with a protein. See Harlow and Lane, *Antibodies, A Laboratory Manual*, Cold Spring Harbor Publications, New York (1988), for a description of immunoassay formats and conditions that can be used to determine selective binding. Examples of types of immunoassays

that can utilize antibodies of the invention are competitive and non-competitive immunoassays in either a direct or indirect format. Examples of such immunoassays are the radioimmunoassay (RIA) and the sandwich (immunometric) assay. Detection of the antigens using the antibodies of the invention can be done utilizing immunoassays that 5 are run in either the forward, reverse, or simultaneous modes, including immunohistochemical assays on physiological samples. Those of skill in the art will know, or can readily discern, other immunoassay formats without undue experimentation.

Antibodies can be bound to many different carriers and used to detect the 10 presence of the target molecules. Examples of well-known carriers include glass, polystyrene, polypropylene, polyethylene, dextran, nylon, amyloses, natural and modified celluloses, polyacrylamides, agaroses and magnetite. The nature of the carrier can be either soluble or insoluble for purposes of the invention. Those skilled in the art will know of other suitable carriers for binding monoclonal antibodies, or will be able to 15 ascertain such using routine experimentation.

The invention also provides a method for serologically diagnosing an EMCR-CoV virus infection of a mammal comprising determining in a sample of said mammal the presence of an antibody specifically directed against an EMCR-CoV virus or component thereof by reacting said sample with a proteinaceous molecule or fragment thereof or an 20 antigen according to the invention

Methods and means provided herein are particularly useful in a diagnostic kit for diagnosing an EMCR-CoV virus infection, be it by virological or serological diagnosis. Such kits or assays may for example comprise a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an antigen and/or an antibody according to the invention.

Use of a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an 25 antigen and/or an antibody according to the invention is also provided for the production of a pharmaceutical composition, for example for the treatment or prevention of EMCR-CoV virus infections and/or for the treatment or prevention of atypical pneumonia, in particular in humans. Preferably a peptide comprising part of the amino acid sequence of the spike protein as depicted in the relevant translations of sequences E and F of figure 3, is used for the preparation of a therapeutic or prophylactic peptide. Also 30 preferably, a protein comprising the amino acid sequence of the spike protein as depicted in the relevant translations of sequences E and F of figure 3, is used for the preparation of a sub-unit vaccine. Furthermore, the nucleocapsid of Coronaviruses, as

depicted in the translation of sequence F, in figure 3, is known to be particularly useful for eliciting cell-mediated immunity against Coronaviruses and can be used for the preparation of a sub-unit vaccine.

5 Attenuation of the virus can be achieved by established methods developed for this purpose, including but not limited to the use of related viruses of other species, serial passages through laboratory animals or/and tissue/cell cultures, serial passages through cell cultures at temperatures below 37°C (cold-adaption), site directed mutagenesis of molecular clones and exchange of genes or gene fragments between related viruses.

10 A pharmaceutical composition comprising a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an antigen and/or an antibody according to the invention can for example be used in a method for the treatment or prevention of an EMCR-CoV virus infection and/or a respiratory illness comprising providing an individual with a pharmaceutical composition according to the invention. This is most useful when said 15 individual comprises a human. Antibodies against EMCR-CoV virus proteins, especially against the spike protein of EMCR-CoV virus, preferably against the amino acid sequence as depicted in translation 2 of sequence E and translation 1 of sequence F in figure 3, are also useful for prophylactic or therapeutic purposes, as passive vaccines. It is known from other coronaviruses that the spike protein is a very strong antigen and 20 that antibodies against spike protein can be used in prophylactic and therapeutic vaccination.

The invention also provides method to obtain an antiviral agent useful in the treatment of atypical pneumonia comprising establishing a cell culture or experimental animal comprising a virus according to the invention, treating said culture or animal 25 with a candidate antiviral agent, and determining the effect of said agent on said virus or its infection of said culture or animal. An example of such an antiviral agent comprises an EMCR-CoV virus-neutralising antibody, or functional component thereof, as provided herein, but antiviral agents of other nature are obtained as well.

30 The invention also provides use of an antiviral agent according to the invention for the preparation of a pharmaceutical composition, in particular for the preparation of a pharmaceutical composition for the treatment of atypical pneumonia, especially when caused by an EMCR-CoV virus infection, and provides a pharmaceutical composition comprising an antiviral agent according to the invention, useful in a method for the treatment or prevention of an EMCR-CoV virus infection or atypical pneumonia,

said method comprising providing an individual with such a pharmaceutical composition.

The invention also comprises an animal model usable for testing of prophylactic and/or therapeutic methods and/or preparations. It is hypothesized that apes can be 5 infected with the EMCR-CoV virus, thereby showing clinical symptoms, and more importantly, similar tissue morphology as found in humans suffering from atypical pneumonia caused by the EMCR-CoV virus. Subjecting apes to a prophylactic or therapeutic treatment either before or during infection with the virus will have a good and useful predictive value for application of such a prophylaxis or therapy in 10 human subjects.

The invention is further explained in the Examples without limiting it thereto.

**Figure legends**

Fig. 1: Phylogenetic relationship for the nucleotide sequences of isolate EMCR-CoV with its closest relatives genetically. Phylogenetic trees were generated by maximum likelihood analyses using 100 bootstraps and 3 jumbles. The scale representing the number of nucleotide changes is shown for each tree. Figure 1a. Maximum likelihood tree of matrix gene nucleotide sequences. Numbers in trees represent bootstrap values. The scale bar roughly reflects 10 % nucleotide differences between related sequences. Figure 1b. Maximum likelihood tree of nucleocapsid gene nucleotide sequences. Numbers in trees represent bootstrap values. The scale bar roughly reflects 10 % nucleotide differences between related sequences.

Fig. 2: Similarity matrix indicating the nucleotide and amino acid identity for the putative Matrix protein (2a and 2b resp.) and for the putative Nucleoprotein (2c and 2d resp.) between the EMCR-CoV virus and closely related coronaviruses. See text for abbreviations.

Fig. 3: Nucleotide sequences from parts of the EMCR-CoV virus. Also included are the putative polypeptide sequences of polypeptides and alignments of the putative polypeptides with that of another member of the Coronoviridae family, where possible (mostly HCoV-229E).

## Examples

### *Specimen collection*

5 Virus was collected from an 8 month old patient suffering from pneumonia using nasal swabs.

### *Virus isolation and culture*

10 Throat swabs were dipped into a culture of tMK cells and passaged four times. Virus was then in Vero-118 cells. One litre of virus containing cell culture supernatant was harvested, and the virus was pelleted in an ultracentrifuge and the virus pellet was resuspended in 1ml PBS.

### *RNA isolation*

15 RNA was isolated from the supernatant of infected cell cultures or sucrose gradient fractions using a High Pure RNA Isolation kit according to instructions from the manufacturer (Roche Diagnostics, Almere, The Netherlands).

### *Sequencing*

20 Purified RNA was sent to BaseClear holding BV (Leiden, The Netherlands) for sequencing.

### *Phylogenetic analyses*

25 Nucleotide sequences were aligned using Clustal W running under BioEdit version 5.0.9. Maximum likelihood trees were created using the Seqboot and DNA-ML packages of Phylipl 5.6 using 100 bootstraps and 3 jumbles. The consensus trees were calculated using the Consense package of phylipl 5.6. These consensus trees were used as usertree in DNA-ML to recalculate the branch lengths from the original sequences.

30 The sequences of EMCR-CoV were compared with those of reference viruses representing each species in the four groups of coronaviruses. These were: human coronavirus 229E (229E), af304460; porcine epidemic diarrhea virus (PEDV) af353511; transmissible gastroenteritis virus (TGEV), aj271965; bovine coronavirus (BoCoV), af220295; murine hepatitis virus (MHV), af201929; avian infectious bronchitis virus (AIBV), m95169, Canine coronavirus (CaCoV), d13096; feline coronavirus (FeCoV),

ay204704; porcine respiratory coronavirus (PRCoV), z24675; human coronavirus OC43 (OC43), m76373, l14643, m933990; porcine haemagglutinating encephalomyelitis virus (HEV), ay078417; rat coronavirus (RtCoV) af 207551) References for the viruses are the numbers of the NCBI catalog (<http://www.ncbi.nlm.nih.gov/entrez/>).

5

In general, coronaviruses, such as EMCR-CoV can be isolated and identified according to the following protocol:

*Specimen collection*

In order to find virus isolates nasopharyngeal aspirates, throat and nasal swabs, 10 bronchoe alveolar lavages, serum and plasma samples, and stools preferably from mammals such as humans, carnivores (dogs, cats, mustellits, seals etc.), horses, ruminants (cattle, sheep, goats etc.), pigs, rabbits, birds (poultry, ostriches, etc) should be examined. From birds cloaca swabs and droppings can be examined as well. Sera should be collected for immunological assays, such as ELISA, molecular-based assays, 15 such as RT-PCR and virus neutralisation assays.

Collected virus specimens may be diluted with 5 ml Dulbecco MEM medium (BioWhittaker, Walkersville, MD) and thoroughly mixed on a vortex mixer for one minute. The suspension is thus centrifuged for ten minutes at 840 x g. The sediment is spread on a multispot slide (Nutacon, Leimuiden, The Netherlands) for 20 immunofluorescence techniques, and the supernatant is used for virus isolation.

*Virus isolation*

For virus isolation Vero-118 cells or tMK cells (RIVM, Bilthoven, The Netherlands) were cultured in 24 well plates containing glass slides (Costar, Cambridge, UK), with the 25 medium described below supplemented with 10% fetal bovine serum (BioWhittaker, Vervier, Belgium). Before inoculation the plates were washed with PBS and supplied with Eagle's MEM with Hanks' salt (ICN, Costa mesa, CA) supplemented with 0.52/liter gram NaHCO<sub>3</sub>, 0.025 M Hepes (Biowhittaker), 2 mM L-glutamine (Biowhittaker), 200 units/liter penicilline, 200 µg/liter streptomycine (Biowhittaker), 1 gram/liter 30 lactalbumine (Sigma-Aldrich, Zwijndrecht, The Netherlands), 2.0 gram/liter D-glucose (Merck, Amsterdam, The Netherlands), 10 gram/liter peptone (Oxoid, Haarlem, The Netherlands) and 0.02% trypsin (Life Technologies, Bethesda, MD). The plates were inoculated with supernatant of the patient samples, 0.2 ml per well in triplicate, followed by centrifuging at 840x g for one hour. After inoculation the plates were

incubated at 37 °C for 1-7 days and cultures were checked daily for CPE. Extensive CPE was generally observed within 5-10 and included detachment of cells from the monolayer..

5 *Virus culture*

Sub-confluent monolayers of tMK cells or Vero clone 118 cells in media as described above were inoculated with supernatants of samples that displayed CPE or with samples taken from a patient.

10 *RNA isolation*

RNA was isolated from the supernatant of infected cell cultures or sucrose gradient fractions using a High Pure RNA Isolation kit according to instructions from the manufacturer (Roche Diagnostics, Almere, The Netherlands). RNA can also be isolated following other procedures known in the field (*Current Protocols in Molecular Biology*).

15

*Sequence analysis*

Sequence analyses were performed as follows: Purified viral RNA (500ng) was converted to cDNA using the SuperScript Choice system (Invitrogen Corp., Carlsbad, CA) by random priming according to the manufacturer's instructions. Blunt-ended, 20 doublestranded cDNA fragments were size-selected on agarose gel to include fragments ranging from 750bp to 4kb. Following purification by spin column (Zymo Research, Orange, CA), cDNA fragments were ligated into pSMART-HCamp (Lucigen Corp., Middleton, WI). The resulting library was electroporated into DH10B ElectroMAX cells (Invitrogen Corp., Carlsbad, CA), and inserts were amplified from individual colonies 25 using pSMART AmpL1 and AmpR1 primers. PCR fragments were sequenced using BigDye 3.1 chemistry and run on a ABI3730 machine (Applied Biosystems, Foster City, CA).

30

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## Claims

(68)

1. An isolated essentially mammalian positive-sense single stranded RNA virus (EMCR-CoV) comprising one or more of the sequences of figure 3.  
5
2. An isolated positive-sense single stranded RNA virus (EMCR-CoV) belonging to the Coronaviruses and identifiable as phylogenetically corresponding thereto by determining a nucleic acid sequence of said virus and testing it in phylogenetic tree analyses wherein maximum likelihood trees are generated using 100 bootstraps and 3  
10 jumbles and finding it to be more closely phylogenetically corresponding to a virus isolate having the sequences as depicted in figure 3 than it is corresponding to a virus isolate of PEDV (porcine epidemic diarrhea virus), HCoV-229E (human coronavirus 229E), PRCoV (porcine respiratory coronavirus), TGEV (transmissible gastroenteritis virus), CaCoV (Canine coronavirus) and FeCoV (feline coronavirus).  
15
3. A virus according to claim 1 or 2 wherein said nucleic acid sequence comprises a open reading frame (ORF) encoding a viral protein of said virus.  
20
4. A virus according to claim 3 wherein said open reading frame is selected from the group of ORFs encoding the viral replicase, nucleocapsid protein, matrix protein or the spike protein.  
25
5. A virus according to claim 1-4 isolatable from a human with respiratory tract disease such as, but not limited to, atypical pneumonia.  
30
6. An isolated or recombinant nucleic acid or EMCR-CoV virus-specific functional fragment thereof obtainable from a virus according to anyone of claims 1 to 5.
7. A vector comprising a nucleic acid according to claim 6.
8. A host cell comprising a nucleic acid according to claim 6 or a vector according to claim 7.

9. An isolated or recombinant proteinaceous molecule or EMCR-CoV virus-specific functional fragment thereof encoded by a nucleic acid according to claim 6.
10. An antigen comprising a proteinaceous molecule or EMCR-CoV virus-specific functional fragment thereof according to claim 9.
11. An antibody specifically directed against an antigen according to claim 10.
12. A method for identifying a viral isolate as an EMCR-CoV virus comprising reacting said viral isolate or a component thereof with an antibody according to claim 11.
13. A method for identifying a viral isolate as an EMCR-CoV virus comprising reacting said viral isolate or a component thereof with a nucleic acid according to claim 6.
14. A method for virologically diagnosing an EMCR-CoV infection of a mammal comprising determining in a sample of said mammal the presence of a viral isolate or component thereof by reacting said sample with a nucleic acid according to claim 6 or an antibody according to claim 11.
15. A method for serologically diagnosing an EMCR-CoV infection of a mammal comprising determining in a sample of said mammal the presence of an antibody specifically directed against an EMCR-CoV virus or component thereof by reacting said sample with a proteinaceous molecule or fragment thereof according to claim 9 or an antigen according to claim 10.
16. A diagnostic kit for diagnosing an EMCR-CoV infection comprising a virus according to anyone of claims 1 to 5, a nucleic acid according to claim 6, a proteinaceous molecule or fragment thereof according to claim 9, an antigen according to claim 10 and/or an antibody according to claim 11.
17. Use of a virus according to any one claims 1 to 5, a nucleic acid according to claim 6, a vector according to claim 7, a host cell according to claim 8, a proteinaceous

molecule or fragment thereof according to claim 9, an antigen according to claim 10, or an antibody according to claim 11 for the production of a pharmaceutical composition.

18. Use according to claim 17 for the production of a pharmaceutical composition for  
5 the treatment or prevention of an EMC-CoV virus infection.

19. Use according to claim 17 or 18 for the production of a pharmaceutical composition for the treatment or prevention of atypical pneumonia.

10 20. A pharmaceutical composition comprising a virus according to any one of claims 1 to 5, a nucleic acid according to claim 6, a vector according to claim 7, a host cell according to claim 8, a proteinaceous molecule or fragment thereof according to claim 9, an antigen according to claim 10, or an antibody according to claim 11.

15 21. A method for the treatment or prevention of an EMC-CoV virus infection comprising providing an individual with a pharmaceutical composition according to claim 20.

22. A method for the treatment or prevention of atypical pneumonia comprising  
20 providing an individual with a pharmaceutical composition according to claim 20.

23. A viral replicase encoded by an RNA sequence comprising the sequences A, B, C, D, E and/or F, or homologues thereof as depicted in figure 3.

25 24. A viral spike protein comprising the amino acid sequence depicted as a translation of (part of) sequences E and F as depicted in figure 3, or a homologue thereof.

25 26. A viral nucleocapsid encoded by an RNA sequence comprising a translation of  
30 (part of) the sequence F as depicted in figure 3 or a homologue thereof.

26. A viral nsp 3 or envelope protein encoded by an RNA sequence comprising a translation of (part of) the sequence F as depicted in figure 3.

27. A nucleic acid sequence which comprises one or more of the sequences A to F as depicted in figure 3 or a nucleic acid sequence which can hybridise with any of these sequences under stringent conditions.

18.11.2003

## Abstract

(68)

The invention relates to the field of virology. The invention provides a new isolated essentially mammalian positive-sense single stranded RNA virus (EMCR-CoV) within the group of coronaviruses and components thereof.

5

68

Figure 1.

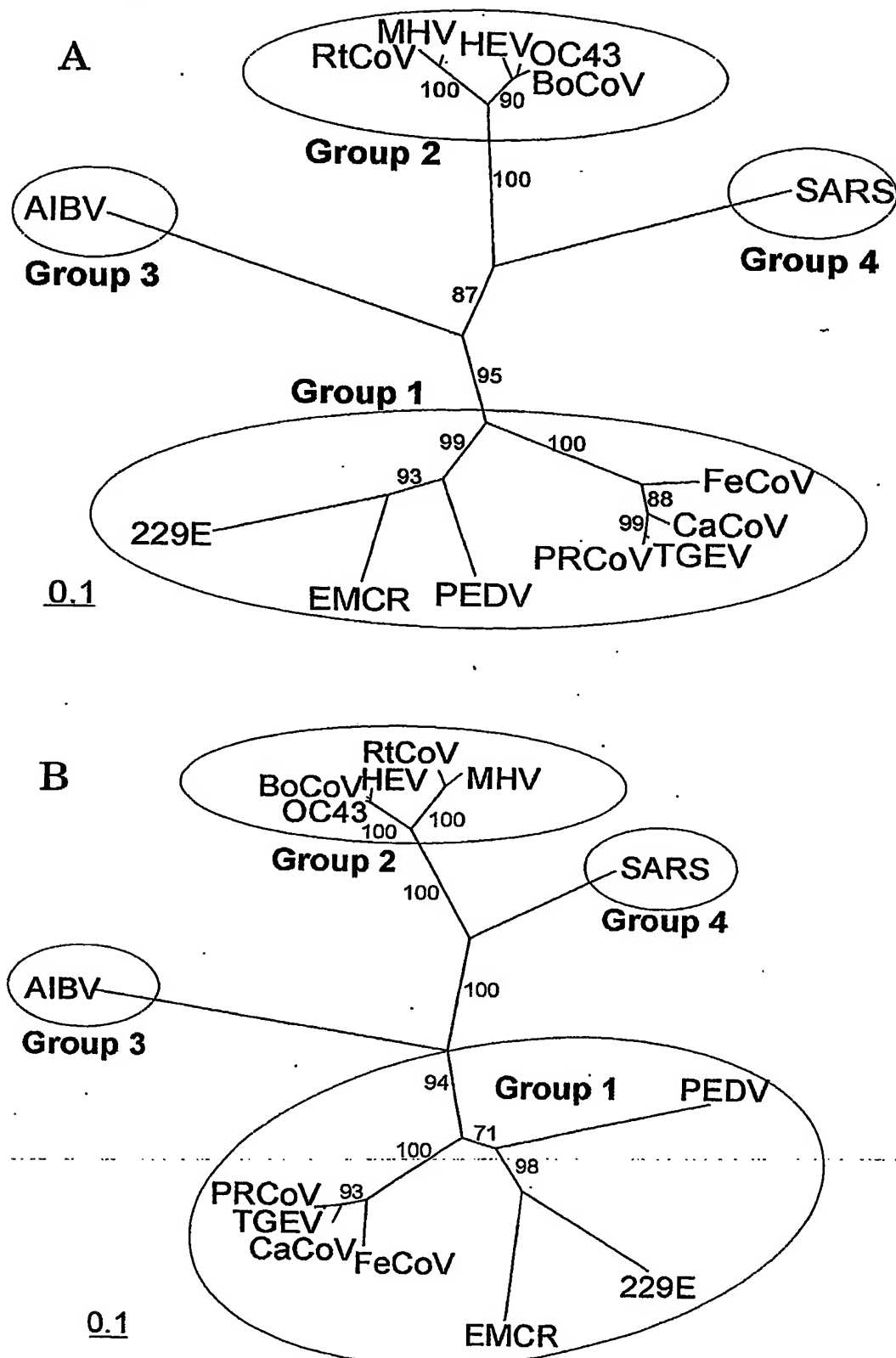


Figure 2a

Figure 2b

Figure 2c

Figure 2d

Seq>	EMCR	229E	PEDV	TGEV	PRCoV	CaCoV	RSDAC	MHV	PHEV	OC43	BoCoV	SARS	AlBV
EMCR	1,000	0,447	0,358	0,336	0,326	0,334	0,344	0,188	0,189	0,179	0,183	0,210	0,173
229E	—	1,000	0,336	0,335	0,304	0,328	0,333	0,196	0,204	0,187	0,190	0,199	0,173
PEDV	—	—	1,000	0,277	0,244	0,272	0,270	0,163	0,168	0,160	0,160	0,184	0,178
TGEV	—	—	—	1,000	0,761	0,963	0,897	0,220	0,223	0,200	0,202	0,232	0,192
PRCoV	—	—	—	—	1,000	0,756	0,763	0,209	0,212	0,185	0,187	0,189	0,185
CaCoV	—	—	—	—	—	1,000	0,879	0,220	0,228	0,202	0,204	0,230	0,192
RSDAC	—	—	—	—	—	—	1,000	0,215	0,221	0,196	0,198	0,216	0,196
MHV	—	—	—	—	—	—	—	1,000	0,894	0,693	0,697	0,285	0,200
PHEV	—	—	—	—	—	—	—	—	1,000	0,680	0,684	0,282	0,208
OC43	—	—	—	—	—	—	—	—	—	1,000	0,948	0,933	0,261
BoCoV	—	—	—	—	—	—	—	—	—	—	0,973	0,261	0,197
SARS	—	—	—	—	—	—	—	—	—	—	—	0,266	0,197
AlBV	—	—	—	—	—	—	—	—	—	—	—	—	0,211
								—	—	—	—	—	1,000

Figure 3

## RNA sequences, implied polypeptides and alignment with one close relative

### 1. Sequence A

### 3762 Nucleotides encoding part of Replicase

ATTCGTTCTATAGAGAATTCTTCTTAACTTGTGCTACTCCTCTCAACTAAAGCAAATTCTAG  
TGCTGTCATTGTTATGGCAGTCCTAGTGTAAATTGAAATTCTGTCAGGTTAGGCAAGTGTG  
ATTTCTGTGTTAACGACTGGTGGTCTGTCCACTAGTGCACACATGATACTTAAGTGTGTTCTGCACTGC  
TTATTGTGGAAGCAACGTTCTGTGTTGGAAACCAATAACTGCTAACATGTTTACAATCAAGTGCACACTTG  
CTGTTGCAAGTGAATCGGAAATTCTCAGGTTTGTGCAATTCTCTGTAGCGCTTCGCTTATAGCGAAG  
CCGCTGACAAGGTTTGTGCAATTCTCTGTAGCGCTTCGCTTACAAGGATTGTGTAACCGGTATTAATGATG  
ACGATTATGTCATTGCAATTGACTGGTACTAATCAGCTTGTGCAAAATTACTTTACTTTCTGATAGACCTCTA  
ATTGCGAGGTTGGCTCATTTCTAACAGCAATTATGTCCTCAGGACTTGTGTTGATGGTAAACCTGTGTTAC  
CAGGAAGTGTGGTTTGTGATAAGTATATGTCGTTTGATGGTAAACATGTG  
AATTAGAGATTACTTAATGATAACTGATAGTATTGTTATTGGTGGTGTCACTTATCAATTAGCATGGGATG  
TTATACGAAAGACCTTCTTATGAACAGCAAATGTTAGCTATTGAGAGCATTCAATTCTGGCACTACAG  
GTCATACTTGAAGTCTGGTGCACAACTCATTAATGCCAACGCCCTAAATATTCTCTAAGGTGTTTGAGTG  
GTGAATGGAATGTCGTGTTAGGCCTTGGCTCACCATTATTACAAATGGTATATCATTGCTAGATATAATG  
TTAAACAGTTCTTAAATGCTTTGTTAAATGCAATTGTCGTTCTGAGAATTGGAGTGTGGTCATGGGATG  
GTTATCATCTTGTGTCGCAACACCTGCTAAGAAACTTGTGTTGTTCTGGTAATGTTGTTCTGGTATG  
TGATCATCACCTCAACTGATGCTGGTGTGTTAAATACTATGCTGGCTTAGTTGTTAAACATATTACTAAC  
TTACTGGTGTGTCATTGCGTGTACAGCTGTTCAATTGATGGAATGTTGGCAACATCTTCTATGATG  
CACTTTGCATAGAAATTCAATTAGACCTTTGCTTGTGTTAACACTTACTTCTAATCAATTACGTCAG  
CTTTCTGGTGTCTGTTACAGAAGATGTTAAATTGCTGCTAGCACTGGTGTATTGACATTAGTCGTGGTA  
TGTGGTCTTACGATGACATATTGACAACAAATAACCTGGTTGTCAGCAAAGCTCTGGGCTTTGATG  
CAATCTGGATGCTTGTGCGCTATTAGCTTGTGCAACTACTACTGGTGGTTGGTTAGGTTGTTAAAGT  
CTATCGCTTCAACTGTTAACTGTTCTAATGGTGTATTATTATGTCGAGATGTTCCAGATGCTTTCAAC  
CAGTTTACCGCACATTACAACAGCTATTGTCGTCATTGATTTCTTAAAGTGTATTAAATTGGTATG  
TTAAATTAAACGACTGGTGAATTGTTCTTACTGAAAATGCTTGTGACTACTGAAGTTGTTG  
GTGTTGCGATGTCGCAATAAGAAAGCCATGTTACTAAAGTAGTTGAGGCTTACAACACTGAAGTTAAGT  
CTGTTATTGAACTTGCACGTGTTATTGCGTCTGTTGATTGCAACCTGTTGACCTGTTGCCCCAAAGGTTAA  
TTGTTATTGCTGGACAAGCTTTCTATAGTGGTTTATGTTTATGTTGATTCTACAACACTGATTAA  
ATGACCCCTGTTTACTGGTGAGTTATTAACTATTAGTTAGTGGTTTAAAGCTTGTGATGGTTTAAACCAT  
AGTTGTTAATGCTAGTCTGCTACAGATGCCATTATTGCTGTTGAGCTGTTGTTATGGTATTAAACTGCA  
TTTTGTTGTCACATGTTGTTGATGGTTGAGTGTCTATTGTTAGACGTTGATCTACATTGCCACACATG  
GTTTAAAGGACTGTTATTGGAGCAATTCTGTCATTGATGTTGAGGCTTGTGTTGACTGATT  
ATAATGCTATTCTGCAAGAGATAAACCTCAATGTGCTATTGTCACCGAGCTAAAGTTGCTTGAGA  
GGTTTACCTAAGTGTGACTGTTGAGTATTGATGATGCCATTGGAATCTTTGTTGAAAAGT  
TTAATTGTTACAGATTGGTAAACACTCTTAAAGCTTACACTTACTCTAATGGTCTTTAGGTAAATTGTC  
AACGTTTACGCTGTTGGTAAATTGCTGTTAATGTTCCATTGTTGAGTATTGTTGTTGAGTGTG  
ACACTGCTGGTGTGTTGCAATTAAATTATGCTGTTAATGTTGAGTATTGTTGTTGAGTGTG  
TAATTGCTAGAGAAAGGTGTGACGTGACTTTCTTGTGTTAGTTGTTGACTTCTTCTATGAAATT  
CGTGTGTTGGTGTAGTAAACCTAATGCCATTGATGTTGAAACATTGATGTTGACTTCTTCTGAGC  
CTAAGGATGGTGGTCAATTGTTGTTCTGATGATTATCTTGTGTTAGGTGAGTGTGACATT  
CATGTAATGGTGTATTGCGCAGTGTGTTTACAAAATTGGCAGGGTAAAATATCTTCTGATGATGTTAG  
TTCATGATGTTGAAACCTACCCATAAAAGTCAGCTCATATTGAGTTGAAGATGATGTTGTTAC  
AGAAGAGTTGGTAAAGTCTATTATTACAGGTGATTGGAAGGTTACATGAAGTTCTTACATGCAATGA  
ATGTCATTGGCAACATATTAAAGTTGCCACAATTATTATGATGATGAAAGAGGGTGTGTTGATGTTCTAAAC  
CAGTTATGATTTCACAATTGGCCTTATTAGTGTGATGATGTTGATGTTGTTGAAACCGAGCACTGATT  
AATTAGAATCTGTTAGAGAAGAGGTGTGATATAATTGAAACAAACCTTGGGAGTGTGAAACATG  
GACAACCTTTCTTTCTTGTGTTAGAGATGAAATTGGGTTGCTGTTAGATGAACTCTGATA  
TTAGTACCAACTTACAGTGTGCAATTCAAAAGCTTTGGGATGATTCTATTGAGATGCAATT  
GTAAAGGTTGATTCAATTGTTCAAAAGTGTGTTAGGTTGCTCATTAAATTGTTGAGTGTG  
AACTCTTGTGAACTTCTTAAAGATAAAATACATGTTCTATAACTTGGGAGATGTGTTGATTG  
AGTTGATGAGCAAGTGTGTTGTTGGATTATGCTTACACAAAACCTTCAAAAGGTGAGAAGCAATT  
CAGCTGTTCTCG

## Putative ORFs

>-out: 140 to 310: Frame 2 57 aa

ASVUVSVFKHWWFCPLVHTLILKWCSTAYCGSNVLSLWKPIТАHVLQSSDTCCCCK

>-out: 267 to 8761: Frame 3 1165 aa

LLTMFYNQVTLAVASDSEISGFGFAIPSVAVRAYSEAAAQGFACRFVAFGLQDCVTGINDDDYVIALTGTNQL  
 AKILLFSDRPLNLRGWLIFSNNSNYVLQDFVVFGHGAGSVVFVDKYMCFGDKPVLPKNMWEFRDYFNDNTDSI  
 IGGVTYQLAWDVIRKDLSYEQQNVLAIESIHYLGGTGTGHTLKGCKLINAQPKYSSKVVLSEWNAVYKAFGSP  
 5 ITNGISLLDIIVKPVFFNAFKCNCSENWSVGAWDGYLSSCCGTPAKKLCVPGNVVPGDVIITSTDAGCGVK  
 YAGLVVKHITNITGVSLWRVTAVHSDGMFVATSSYDALLHRNSLDPFCFDVNTLLSNQLRLAFLGASVTEGVK  
 ASTGVIDISAGMFGLYDDILTNNKPFVFKASGLFDIAIWDAFVAAIKLVPTTGLVRFVKSIASTVLTGSNGV  
 IMCADVPAFQPVYRTFTQAICAAQDFSLDVFKIGDVFKRGLGDYVLTENALVRLTEVVRGVRDARIKKAMFT  
 VVVGPTTEVKFSVIELATVNLRLVDCAPVCPKGKIVVVIAGQAFFYSGGFYRFMVDSTTVLNDPVTGELFYTI  
 10 FSGFKLDGFNQFVNASSATDAAIAVEVLLSDFKTAVFVYTCVVDGCSVIVRRAFATHVCFKDCYSIWEQFC  
 DNCGEPWFLTDYNAIQLQSNPQCAIVQASESKVLERFLPKCPEVLLSIDDGHLWNLFVEKFNFTDWLKL  
 LTSNGLLGNCAKRFRRLVVKLLDVYNGFLETVCVVHTAGVCVYAVNVPVYVVISGFVSRVIRRERCDVTFPC  
 SCVTFFYEFLLTCFGVSKPNAIDVEHLELKETVVFEPKDGGQFFVSDDYLWYVVDDIYYPASCNGLVPAFTKL  
 GKGKISFSDDVIVHDVEPTHVKLIFEFEVDVVTSLCKKSFGKSIYTGDWEGLHEVLTSAINVIGQHILKLPQFY  
 15 YDEEGGYDVSKPVMISQWPISDDSDGCVVEASTDFHQLESVREEDVIEQPFGEVEHALSIRQPFSSFRDELG  
 RVLDQSDNNCWISTTLIQLQLTKLDDSIEMQLFKVGKVDSIVQKCYELSHLISGSLGDSGKLLSELLKDKYTC  
 ITFEMSCDCGKKFDEQVGCLFWIMPYTKFKVVRTNSAVL  
 >-out: 472 to 738: Frame 1 89 aa  
 LVLLISFVPKFYFFLIDLLICEVGSFFLTAIMFFRFLMLFLAMVQEWFVLWISICVVLMVNLCYLTCGNLEITL  
 IIILIVLLVVSLIN  
 20 >-out: 973 to 1125: Frame 1 51 aa  
 LLNQFSLMLLLNAIVVLRIGVLVHGMVIYLLVVAHLLRNFLVFLVMLFLVM  
 >-out: 2026 to 2316: Frame 1 97 aa  
 MTLFLLVSYFILLSVVLSMVLTISLLMLVLLQMPILLSCCCYRILKQFLCTHWWLMVVVSLLDVMLHSPHM  
 VLRTVIVFGNSNALSIIIVVSHGF  
 25

### Alignment

>gi|281286|pir||S28600 hypothetical protein 1a - human coronavirus  
 gi|59491|emb|CAA49877.1| ORF1a [Human coronavirus 229E]  
 30 Length = 4085  
 Score = 882 bits (2280), Expect = 0.0  
 Identities = 470/1159 (40%), Positives = 675/1159 (58%), Gaps = 7/1159 (0%)  
 Frame = +3

35 Query: 276 MFYNQVTLAVASDSEISGFGFAIPSVAVRAYSEAAAQGFACRFVAFGLQDCVTGINDDY 455  
 M N+VTLAVASDSEIS G + + AVR YSEAA+ GF+ACRFV+ LQDC+ GI DD  
 Sbjct: 1 MACNRVTLAVASDSEISANGCSTIAQAVRRYSEASNGFRACRFVSLDLQDCIVGIADDT 60

40 Query: 456 YVIALTGTNQLCAKILLFSDRPLNLRGWLIFSNNSNYVLQDFVVFG-HGAGSVVFVDKY 632  
 XV+ L G L I+ FSDRP L GWL+FSNSNY+L++FDVVFG G G+V + D+Y+  
 Sbjct: 61 YVMLGLHGNQTLFCNIMKFSDRPFLHGWLVPNSNSYLLLEFDVVFKGRRGGNVTYTDQYL 120

45 Query: 633 CGFDGKPVLPKNMWEFRDYFNDNTDSIVIGGVTYQLAWDVIRKDLSYEQQNVLAIESIHY 812  
 CG DGKPV+ +++W+F D+F +N + I+I G TY AW RK L Y++QN LAIE I Y  
 Sbjct: 121 CGADGKPVMSEDWLWQFVDHFGEN-EEIIINGHTYVCAWLTKRKPLDYKRQNNLATEIEY 179

50 Query: 813 L-GTTGHTLKGCKLINAQPKYSSKVVLSEWNAVYKAFGSPFITNGISLLDIIVKPVF 989  
 + G HTL++G L AK K SSKVVL + +YK FGSP +TNG ++L+ KPVF  
 Sbjct: 180 VHGDALHTLRNGSVLEMAKEVKTSSKVVLSDALDKLYKVFSPVMTNGSNILEAFTKPVF 239

55 Query: 990 FNAFKCNCSENWSVGAWDGYLSSCCGTPAKKLCVPGNVVPGDVIITSTDAGCGVKYY 1169  
 +A V+C CG++WSVG W G+ SSCC + KLCVPGNV PGD +IT+ AG G+KY+  
 Sbjct: 240 ISALVQCTCGTKSWSVGDWTGFKSSCCNVISNKLCVPGNVKPGDAVITQQAGAGIKYF 299

60 Query: 1170 AGLVVVKHITNITGVSLWRVTAVHSDGMFVATSSYDALLHRNSLDPFCFDVNTLLSNQLRL 1349  
 Gt. +K + NT GVS+WRV A+ S... FVA+S++ ... H. N. +D. FCF+V- +++++ RL  
 Sbjct: 300 CGMTLKFVANIEGVSVWRVIALQSVDCFVASSTFVEEEHVNRMDTFCFNRNSVTDECRL 359

65 Query: 1350 AFLGASVTEVDVKAFASTGVVIDISAGMFGLYDDILTNNKPFVFKASGLFDIAIWDAFVAAI 1529  
 A LGA +T +V+ ++GVIDIS G F +YDDI +KPWFVKA +F W A +A+  
 Sbjct: 360 AMLGAEMTSNVRQVAVGVIDISTGWFVDVYDDIFAEASKPWFVKAEDIFGPCWSALASAL 419

70 Query: 1530 KLVPTTGGLVRFVKSIASTVLTGSNGVITIMCADVPAFQPVYRTFTQAICAAQDFSLDV 1709  
 K + TTG LVRFVKS + + + V G I + A VP+ F + F AI FD +++  
 Sbjct: 420 KQLKVTGELVRFVKSICNSAVVGGTIQILASVPEKFLNAFDVFTAIQTVFDCAVET 479

Query: 1710 FKIGDVFKFRLGDYVLTENALVRLTEVVRGVRDARIKKAMFTKVVVGPTTEVKFSVIEL 1889  
 I F ++ DYVL +NALV+L T ++GVR+ + K + VVVG T EVK S +E  
 Sbjct: 480 CTIAGKAFDKVFDYVLLDNALVKLVTKLKGVRERGLNKVKYATVVVGSTEEVKSSRVER 539

Fig 3. (cont)

5	Query: 1890	ATVNLLRLVDCAPVVCPKGKIVVIAGQAFYSGGFYRPMVDSTTVLNDPVFTGELFYTIKF +T L + + + +G VVI A+F S G++R M +VL V+ + +	2069
	Sbjct: 540	STAVLTIANNYSKLFDEGYTVVIGDVAYFVSDGYFRLMASPNSVLTTAVYKPLFAFNVN 599	
	Query: 2070	SGFKLIDGFNHNQFVNASSATDAI I A V E L L I L S D F K T A V F V Y T C V V D G C S V I V R R D A T F A T H V G + + F V + A++ V + + + F + Y+ V + IV+ + + +	2249
	Sbjct: 600	MGTRPEKF-PTTVTCENLESAVLFVNNDKITEFQ--LDYSIDVIDNEIIVKPNISLCVPL 655	
10	Query: 2250	CFKDCYSIWEQFCIDNCGEPWFLTDYNAILQSNNPQCAIVQASESKVLLERFLPKCPEVL +D W+ FC E WF DY A + + A V+A+ESK ++ +P CP +L	2429
	Sbjct: 656	YVRDYVDKWDDFCRQYSNESWFEDDYRAFISVLDITDAAVKAAESKAFAVDTIVPPCPSIL 715	
15	Query: 2430	LSIDDGHLWNLFVEKFNFTWDXXXXXXXXXXXXCAKRFRRVLVKLLDVYNGFLET ID G +WN ++ N V DW CAKRF+R L LL+ YN FL+T	2609
	Sbjct: 716	KVIDGGKLIWNGVIKVNNSVRDWLKS LKLNLTQQGLLGTCAKRFKRWLGILLEAYNAFLDT 775	
	Query: 2610	VCSVVHTAGVC1KYYAVNPYVVISGFVSRVIRRERCD--VTFPCVSCVTFFYEFLLTCF V S V G+ K YA + PY+VI V +V + + FP + F F F	2783
20	Sbjct: 776	VVSTVKIGGLTFTKYAFDKPYIVIRDIVCKVENKTEAEWIELFPHNDRIKSFSTFESAYM 835	
	Query: 2784	GVSKPNAIDVHEHLELKETVFEVKDGGQFFVSSDDYLWYVVDDIYYPASCNGVLPVAFTKL ++ P D+E +EL + FVEP GG V D++++Y D +YYP++ +LPVAFTK	2963
	Sbjct: 836	PIADPTHFDIEEVELLDAEFPVEPGCGGILAVIDEHVFYKKDGVVYPSNGTNILPVAFTKA 895	
25	Query: 2964	AGGKISFSDDVIVHDVEPTHKVKLIFEFEDEDVVTS LCKKSFGKSIIYTGDWEGLHENVLTS AGGK+SFSDDV V D+EP + +VKL FEFED+ + +C+K+ GK I + GDW+ + + S	3143
	Sbjct: 896	AGGKVSFSDDVEVKDIEPVYRVKLCFEFEDEKLVDCEKAIGKKIKHEGDWDSFCKTIQS 955	
30	Query: 3144	AMNVIGQHIKLPQFYIYDEEGGYDVSKPVMISQWPIS---DDSDGCVVEASTDFHQLESV A++V+ ++ LP +YIYDEEGG D+S PVMIS+WPS + + + + D + + V	3314
	Sbjct: 956	ALSVVSCYVNLPTYYIYDEEGGNDLSSLPMVISEWPLSVQQAQQEATLPDIAEDV--VDQV 1013	
35	Query: 3315	REEVDIIIEQPFGVEHALSIROQFSFSRDELGVVRVLDQSDNNCWCISXXXXXXXXXXD E I + +V+H +S PF F + G++L Q DNNCW++ D	3494
	Sbjct: 1014	EEVNSIFDIETVDVVKHDVS---PFEMPFEELNLKILKQLDNNCWVNSVMLQIQLTGILD 1070	
	Query: 3495	DSIEMQLFKVGKVD SIVQKCYELSHLXXXXXXXXXXXXXXYTC SITFEMSCDCGK MQ FK+G+V + + + +CY I +T + + C C	3674
40	Sbjct: 1071	GDYAMQFFKMGRVAKMIERCYTAEQCIRGAMGDVGLCMYRLLKDLHTGFMVMDYKCSCTS 1130	
	Query: 3675	KFDEQVGCLFWIMPYTKLF 3731	
		E+ G + + P K F	
	Sbjct: 1131	GRLEESGAVLFCTPTKKAF 1149	

## 2. Sequence B

50 1610 nucleotides encodes part of replicase  
TTCTGCCTATGGAGGTAGGTATGATTAAATGGTCAGTATTGAGCGATATCTAGAGAATTCTGCTGAAAATGG  
TATTCCACTTATGCCCTTCTTAGTTGTGGTATTTGGTGAAGGATTGAAAATTCTCTAAAGCTTTGTTAG  
TTGTGACATTAATAAACCATTGCAGTTTATTCTCAAATGAAGAACAGCTGTTCTTAAGTTTTAGA  
TGGTTTAGATTTAACACCAGTCATTGACGATGTTGATGTTAAACCTTTAGAGTTGAAGGTAATTTCATT  
CTTGATTGTGGTGTCAATGCCCTGGATGGTGAATTACTTATTACTAACTCTATTAAATGTTGGATAAA  
55 ACAAGGACAATTATGGACACAAAACCTTAATGGTATTGCAACAGGCAGTCTGATTATCTGCTACAGTTAA  
AACTGTACAGCTGGTAATTGGTTAAACCTTGTGGTGGAGAGTTGACCAATTATGTTGTTGACCATCGAT  
AAATGATCTTCTTGTAAAAACCTTGGTGTGGTGTGCGTAAACCTTAAATGAGATTGAAAACCTTGTGTTATTG  
CAATGTCCTGCATTGATGTTGAAAAAGCTTCTTCAGTTGACTTTAACTGTTAAATTGTTGAGAGAG  
TAATGTTATGGATGTTAACGACTGTTTAAGAATGATAATGTTGAGTTGAAAATTACTGAAGAGTGGTATTAAATGT  
60 TAAAGATGTTGTTGAGCTCTCTAACGACTGTTTAAGAATGATAATGTTGAGTTGAAAATTACTGAAGAGTGGTATTAAATGT  
TGAAGGTGTTTACCTATTAATACTGATACTGTTCTATCTGTAGCTCCAGAACAGTTGACTGGTTGCTTTTACGG  
TTTGAAAAGGCACTTTTGCTTGGATGTAAGCCATATGGTTACCCATAATGATTGTTGGGTT  
TAGAGTTCTGGGACCAACGACAATAATTGTTGGGTTAATGCAACCTTGTATAATTACAGTATCTAACGCTAC  
65 TTTAAATCTAACGGGTTAAATGTTCTTGGAACAAATTGTTACAGGGTGTGACAGGTTGACCTTTGTTAGTTTAT  
TTATTTTATAACTATGTCCTAACAGGGTCAAAAGGGTGTGACAGGTTGACAGGCTTACAGACTTTGTTAGTTTAT  
GATTAGTGTATTCTTGTACTCTTGAACAAATTCACACTTGTGACATTGTTGAAAAGTACTGTTGAGTTAA  
AAGTGTGTTGCTGTGCTAGTGTGCTTAAAGATGGTGTGATGTTGGTTTGTCCACACAGACATAATTGCG  
70 TTACAGTGTAAAGTTGTTAACGGACGTGTTATTACAAATTTCAAGGTTCTTGTGATAACGGTCACTATGTTGATGCTGC  
GTTGCTTAAGGTTGCTTATACACATTTCAGGTTCTTGTGATAACGGTCACTATGTTGATGCTGC  
TAATAATGCTGTCTATGATGGTGTCTGTTATTGCTTCAGATTGCTACTTTAGCTGTTACAGCTATTGTTG  
AGTAGGGTGGTTGTTGTAACATCTAACCCACAAACG

Putative ORfs

>-out: 32 to 1609: Frame 2 526 aa  
 MVS IERYLENSSENGIPLMPLLSCGIFGVRIENS LKALFSCDINKPLQVFVYSSNEEQAVLKFLDGDDLT PVID  
 5 VDVVKPFRVEGNFSFFDCGVNALDGDYIYLLFTNSILMDKQGQLLDTKLNGILQQAVLDYLATVKTVPAGNLVK  
 VVESCTIYMCVVP SINDL SFDKNLGRCVRLNRLKTCVIANVPAIDVILKLLSSILTVKVVESNVMDVNDCF  
 NDNVVLKITEDGINVKD VVVESSKSLGKQLGVVSDGVDSFEGVL PINTDTVL SVAPEVDWVAFYGFEKAALFAS  
 DVKPYGYPNDFVGGFRVLGTTDNNCWVNATCII LQYLKPTFKSKGLNVLWNK FVTGDVGPVFSFIYFITMSSKG  
 KGDAAEALSKLSEYLISDSIVTLEQYSTCDICKSTVVEVKSAVVCASV LKDGC DVGFCPHRK LRSRVKFVN GR  
 10 VITNVGEPIISQPSKLLNGIA YTTFSGSFDNGHVVYDAANNAVYD GARLFASDLSTLAVTAIVVVGGCVTSNF  
 N  
 >-out: 366 to 524: Frame 3 53 aa  
 CWINKDNYWTQNL MVFCNRQFLIILLQLKLYQLVIWLNL LRVV PFICVLYHR

Alignment

15 >gi|12175747|ref|NP\_073549.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 gi|30179827|sp|Q05002|R1AB\_CVH22 Replicase polyprotein 1ab (pp1ab) (ORF1ab polyprotein) [Includes:  
 Replicase polyprotein 1a (pp1a) (ORF1a)] [Contains: p9;  
 p87; p195 (Papain-like proteinases 1/2)  
 (PL1-PRO/PL2-PRO); Peptide HD2; 3C-like proteinase  
 (3CL-PRO) (3CLp) (M-PRO) (p84); Unknown protein 1; p5;  
 p23; p12; Growth factor-like peptide (GFL) (p16);  
 RNA-directed RNA polymerase (RdRp) (Pol) (p100); Helicase  
 (Hel) (p66) (p66-HEL); Unknown protein 2; p41; Unknown  
 protein 3]  
 20 25 >gi|12082740|gb|AAG48591.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 Length = 6758

Score = 429 bits (1104), Expect = e-119  
 Identities = 233/585 (43%), Positives = 323/535 (60%), Gaps = 18/535 (3%)  
 30 Frame = +2

Query: 41 IERYLENSSENGIPLMPLLSCGIFGVRIENS LKALFSCDINKPLQVFVYSSNEEQAVLKFLDGDDLT PVID 220  
 I+ Y ++E G PL P+LSCGIFG++E SL+ L K ++VFVY+ E V F  
 35 Sbjct: 1372 IKAYNTINNEQGTPPLTPI LSCGIFGIKLETSLEVLLDVCNTKEVKVFVYTDTEVCKVKDF 1431  
 Query: 221 LDGLDLTPVIDD-----VDVVK----PFRVEGNFSFFDCGVNAL-DGDYIYLLFTNSIL 364  
 + GL ++ V V+K P+RV+G FS+F + + D +LFT+S+L  
 Sbjct: 1432 VSGLVN VQKVEQPKIEPKPVSVIKVAPKPYRVDGKFSYFTEDLLCVADDKPLIVLFTDSML 1491  
 40 Query: 365 MLDKQGQLLDTKLNGILQQAVLDYLATVKTVPAGNLVKLVVESCTIYMCVVP SINDL SFD 544  
 LD +G LD L+G+L A+ D + K +P+GNL+K + S +YMCVVP S D D  
 Sbjct: 1492 TLDDRGLALDNALSGVLSAAIKDCVDINKAIPSGNLIKFDIGSVVVYMCVVPSEKD KHL D 1551  
 45 Query: 545 KNLGRCVRKLNRLKTCVIANVPAIDXXXXXX FV VESNVMDVNDCFKNDNVVL 724  
 N+ RC RKL NRL ++ +PA FV E + + +  
 Sbjct: 1552 NNVQRCTRKL NRMCDIVCTIPADYI LPLV LSSLT C N VSFVGELKAAEAKV-----ITI 1605  
 Query: 725 KITEDGINVKD VVVESSKSLGKQLGVVSDGVDSFEGVL PINTDTVL SVAPEVDWVAFY 898  
 K+TEDG+NV DV V + KS +O+GV+ +D G +P +NT +L+ A +VDWV FY  
 50 Sbjct: 1606 KVTEDGVNVHDVTVT DKSFEQQVGVIADKDKDLSGAVPSD LNTSELLTKAIDVDWVEFY 1665  
 Query: 899 GFEKAALFASLDV KPYGYPNDFVGGFRVLGTTDNNCWVNATCII LQYLKPTFKSKGLNVL 1078  
 GF+ A FA++ +D + Y + V G RVL T+DNNCWVN A CI LQY KP F S+GL+  
 55 Sbjct: 1666 GFKDAVTFA T D HSAFAYE SAVVNGIRVLKTSDNNCWVN A VCIALQYSKPHFISQGLDAA 1725  
 Query: 1079 WNKFV TGDVGPFVSEIYFITMSSKGQKGDAEEALSKLSEYLISDSIVTLEQYSTCDIC 1252  
 WNKFV GDV FV+F+Y++ KG KGDAE+ L+KLS+YL +++ V LE YS+C C  
 Sbjct: 1726 WNKFV L GDV E I FVAFV Y Y V A RLMKGDKGDAE D T L K L SKYLANEAQVQLEHYSSC V E C D A 1785  
 60 Query: 1253 --KSTVVEVKSAVVCASV LKDGC DVGFCPHRK LRSRVK FVN GRV V ITNVGEPIISQPSK 1426  
 K++V + SA+VCASV +DG VG+C H K SRV+ V GR +I. +V + S+  
 Sbjct: 1786 KFKNSVASINSAI VCA SV KRDGVQVG YCVHGI KYY SRV SRV GR AII V S V E Q L E P C A Q S R 1845  
 65 Query: 1427 LLNGIA YTTFSGSFDNGHVVYDAANNAVYD GARLFASDLSTLAVTAIVVVGGCV 1591  
 LL+G+AYT FSG D GH Y VYD A ++YDG R DLS L+VT++V+VGG V  
 Sbjct: 1846 LLSGVAYTAFSGPVDKGHYTVYDTAKKSMYDGDRFVKHDL SLLS VTS VVMVGGYV 1900

**3. Sequence C**

6017 nucleotides; Encodes part of Replicase

CGAGAACAGCTGATTGTTATTTGTTAATTTGTTAAACGTATTCGCGTTGGACTT  
 5 TTATATTTGTTGACAATTATAGTACTTTGGCTCTTCTAGGCTTCATCAGAACAGTGGTTTACAT  
 TTTGTCGCGTTGATGTTTATGTAATGAGTTTCTAGCTaCATTATTGTCGAAAATGTTTATTTGTTAGA  
 CATATTATTGTTGGCTGTAATAATGCTGACTGTTAGCTGTTCTAAAAGTGTAGACTAAACGTGTAACCTT  
 CAAACTATTATTAAATGGTATGCAAAATCATTCTATGTTAATGCTAATGGTGTACTTGTGTTCTGTAATAAACAT  
 AACTCTTTGTTAATGTTGATTCTTTGGGCTGGTAACTTTTATTAAATGGTGTATTGCAAGAGAGCTT  
 10 GGTAATGTTGTTAAAACAGCTGTTCAACCCACAGCTCCGATATGTTATTGATAAGGTAGATTGTTAAT  
 GGATTATTATCGCTTATAGTGGTGTGACACTTTGGCGGTATGACTTTGACATTACTGAATCTAAGTATAGTGT  
 AAAGAGGTTCTGAAGAAATTGTAATGTTAGAAAATTTTATTGTTACAATAATAGTGGTAGTAACATTACACAG  
 ATTAAAATGCTTGTGTTATTCTCAATTGTTGTTGAACTTAAAGTGGTAAATTCAAGGTTGTTGTA  
 15 ACTTTATCAGTGATTAAATGGTGTGTTGATAAGGCATATGTTGATGTTGTTGTAATAGTTTTAAGGAG  
 CTAACGTGTAACATGTCATGGCTGAAATGTAAGCTACACTGGTTGACTGTTCTGATGATGATTGTTCA  
 GCTGTTGCAATGCAATAGGTATGACGTTGCTTCAGATTGTCATTAAATAATTTTTATTCTTATGCT  
 AAACCTGAAGATAAGTTGTCGTTATGACATTGCTTGTATGCGTGCCTAAGGTTGTTAACCATAAT  
 20 GTTTTAATCAAAGAGTCATAACCTATTGTTGGGCTGTCAGGACTTTAATAACTCTTCTCAAGAACGTAAGAAG  
 TACCTGTTAAAACAACAAAGCAAGGGTTGACTTTTATTAAATGATAACCAAGCAATTACACAA  
 GTCCTGCTACTAGTATAGTGGCAAAACAGGGTGCTGGTTAAACGTTACTTAAATTCTGTTGATGTTATGTT  
 TTATTGTTGTCATTGTTATTGTTGTCATTATTGATTATAACAAACACTGTAACAGTCTTATGTT  
 GATTAAAGTACATTGAGAATGGTCAGTTGAAAGGTTGTTGACCTTACACTGTTGCGTAATGTTTGAT  
 25 AATTAAATCAATGCCATGAGGCTAAGTTGGTGTGTTACTACTAATAGTGTAAATGCTCTATAGTGTGTT  
 GTTTCAGAGCGTATTAAATGTTGTTCTGGTGTCAACAAATGTTATTGTTAGGAAAGACTCTGTTTACA  
 TTACAGGCTGCTTTGGAAACACAGGTGTTATGACTTGTGTTACACTAGTGTAAAGTGTATT  
 30 AATTCTGCTTGACTAGGTTGGAAGGTTGGGTGACAATGTTATTGTTACAACACTGATCTTATTGAAAGGT  
 TCTAAACCTTATAGTATTTCACGCCAATGCTTATTAAAGTGTAAATGCTACGTTTCCAGAA  
 ATTTCAGCTAGAGGTTGGCTTACGTTACTATTAGAACTTGGCTCACGTTATTGTTAGGTTGGAATGCCGT  
 GACTCACAAAGGTGTTGGTTGGTGTAAATGGTGTATTGTTAGTGTAAATGTTGACGTGTTGATGACGGTTACATT  
 35 TGTGTTGATGGCTTATAGACCTCTGTTAAATGTTACTCTCAATCTTGTGTTAGTGTAAAGGTTGCTATGTT  
 TCTGGACATATGTTAAATTCTTTGCAKCATTTATTACATTGTTGCTTTAGTTACTAAATTAAA  
 CGTGTGTTGGTGTCTTCTTATGGTGTGTTACTGTTGTTGCAACTTGTGTTAAATAACATTCTTATGTT  
 GTTACTCAAATTATTATTGTTATGTTGCTTATGCTATTGTTGTTACTAGGACAGTGCCTTATGCT  
 TGGATTGGCATATTGCAACATTGCTTGCATACTTCTGTTAAATACCATTGGGGCTTCACATGGTTAGTT  
 GCTGCAATTAGGCTTACCTAAATGTTAAAGTTAAAATCTCTACTCAATTGTTGAAAGGTGATAAGGTT  
 40 ATAGGTACTTTGAGAGTGTGCTGTCAGGTCACATTGTTCTTGACATGCTTCTTAAAGGCTGATAAAACT  
 ATTTCACCTGAGAAATTGCTGCAAGTTATAAAATATAAAATTATAGTGGTAGTGTGACTGAG  
 GCTGATTATGTTGCTGTTATGCTCATTAGCCAAGGCTATGTTAGATTACGCAAAGATCATAATGACATG  
 TTATATTCTCACCTACCAATTAGCTACAATTCCACCTTACAATCTGGCTTAAAGAAGATGGCACAACCATTGGT  
 TGTGTTGAGAGATGTGTTGGCTGCTGTTATGGTAGTACTGTGCTTAATGGAGTTGGTTAGGTGACACTGTT  
 ACTTGTCCCTAGACATGTCATAGCACCATCAACCAACTGTTCTTATTGATTATGATCATGCAATAGTACTATGCGT  
 TTGCAATAATTTCAGTGTCTCATAATGGTGTCTTCTGGGAGTTGTTGTTACAATGCACTGGTTCTGTTG  
 CGTATTAAAGGTTACAATCTAATGTCATAACACCTAAACATGTTTAAACGTTGAAACACTGGCTCTCTTT  
 AATATTTCAGCTGTTATGAAAGGTTGCTGTTCTGGTTATAATGTTAGAAATGATGGTACTGTTGAGTTGTT  
 45 TTACACCAAATTGAGTTAGGTAGTGGTGTCTGTTACTGGTAGTGTGTTATGGTAATTGGT  
 GACCAACCTAGTTGCAAGTTGAGAGTGCAACCTTATGCTATCAGATAATGTTGTTGCTTGGTTATGCTGCT  
 TTGTTGAATGGTTGAGGTGGTGGTGTGCTCAACTAGAGTTAATGTTGATGGTTAAATGAATGGCTATGGCT  
 AATGGTTATAACAATTGTTCTAGTGTGAGTGCTATTCTATTGTTGCAAGGTTGGTAAACACTGGTTAGTGTGAAACAA  
 TTGTTAGCTCCATTCAACATCTTCTGTTATAATGTTAGAAATGATGGTACTGTTGAGTTGTT  
 50 GAGTTCACACTAGCTGAAGTTGTAAGCAGATGTGTTAAACTGTTGAAAGTGGTATTGGTT  
 AAAAACATGTTTATTAGCTGTTCTTCTCACAATGTTGGGAGCAACTCTTAAACACTATATGG  
 ATAAACCTGTTATAACTTACACCTATATTGTTACTTTGTTGTTACTGTTGTTAAATGTTTCTTAA  
 CATAAGTTTTGTTGCAAGTATTGTTACTCTGTTATTGCAACTGCTTATATAATTGTTTGGAT  
 TATTACATAGTAAAATTGGCTGACCATTAAACTATAATGTTCAGTATTACAAATGGATGTCAGGGTTA  
 55 GTTAATGTTGGCTGTTATTGTTGTTACACACTTATTGTTATAGTGGTGTACTTTGAGTTGCTTGGTT  
 TTTACATATGTTGTTCTTATAGCAGTGTCTTACACTTATTGTTATAGTGGTGTACTTTGAGTTGCTTGGTT  
 ATGTTTATTGTCATACTAGTGTGTTACATTGGTGTACATTGGTGTCTTACGTTGATTATATT  
 TTTTCACCTGTTAAAGTGTATTAGTGTGTTGTTGTTGTTGTTGTTAAATGTTACTATGGTTATGTT  
 TTAGTTGTTGACTATTGGGGATTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTT  
 60 AAGGTGAGTGTGCTGTAATTAAACATGGTGTCTGTTGCTAATGGACTTCATGCACCATATGACCTTGTGACT  
 TGTTTATCATTCAAATTACTGGTATTGGTGGTGTGACCGTTGTTAAACAAATTCAACTGTCACATCCAAACTGACT  
 GATTGAGAGTGTACTAATGTTGTTATTGGGTTGTTGCTAGTATGAAACATTGCACTGGTGTACT  
 GCTTATTGTTGTTGTTACACAATAAGGTTAATCTTGTGATGACCCAGAAAAAGCTCAAGGTATGTTGTTAGCA  
 CTCCCTGCGTTCTTCTAAGTAAACATAGTGTATTGTTGTTGATGGCTTATTGATTCTTATTTGATAATAGT  
 65 AGCACCCCTGCAAGAGTGTGTTGCTCATCATTGTTAGTATGCCATCATATAATTGCTTATGAAAATGCTAGACAAGCT  
 TATGAGGATGCTATTGCTAATGGATCTCTCTCAACTTAAACAAATTGAAAGCGTGCCTGATGAAACAGCTGCTACTCAGATGTAT  
 AAAGAACCGCTCTGTTAATAGAAAATCTAAAGTTAGTGTGCTATGCACTCTTACTTTGGAAATGTTAAGA

CGTTGGATATGTCTAGTGTGAAACTGTTTGAAATTAGCACGTGATGGTGTGCCCCATTGTCAGTTATACC  
GCAACTTCAGCTTCCAAACTAACTATTGTTAGTCCAGATCTTGAATCTATTCTAAGATTGTTGTGATGGTTC  
GTTCAATTATGCTGGAGTTGGACACTTAATGATGTTAAAGACAATGATGGTAGACCTGTTCATGTTAAAGA  
ATTACAAGGGAGAATGTTGAAACTTGCATGGCCTCTTATCCTTAATTGTAACGTGTTAAACTCTAAAAA  
5 AATGAAATTATGCCTGGTAAACTTAAGCAAAACCTATGAAAGCTGAGGGTGTGATGGTGGTGTGTTAGGTGATGG  
AATGCTTTGTATAATACTGAGGGTGGTAAACTTTATGTTATGCTTATATTCTAATAAAGCTGACCTTAAATT  
GTTAAGTGGGAGTATGAGGGTGGTGTGCAACACAATGAGTTAGACTCTCCTTGTGCGATTATGGTCGAAACACC  
AATGGTCCTCAAGTGAAGTATTGTTGTTAAATTAAACCTTACGTAGAGGGGCCGTTCTGGTTT  
10 ATAGGTGCCCCACAATTGCTCTAACAGCTGTAACAAACTGAAATTGGCTGTTAATTCTGGACTTTAACGTCTTG  
GCTTTTCTGTTGATCCAGCAACCACCTACTTGGAGCTGTTAACATGGTGCACAAACCTGTAAGTAATTGTAT  
AAGATGTTATCTAATGGTGTGGTAAATGGTCAAGCTATAACAACACTAGTGTAGATGCTAACACCAATCAAGATT  
TATGGTGGAGCGTCTATTGTTGTTGCTGGGCCCCACGTTCTCACCCCTAGTATGGATGGTTACTGTAAGTT  
15 AAGGGTAAATGTTGCTAGGTTCTATTGGTGTGGGATCTATTAGGTTTGTGTTAGAAAATAATGTTGTTAAG  
GTTTGTGGGTGTTGGGACACGGGTGTGCTTGTGATCGTACAACCAATTCAAAGTGTGACATTCTTATTAA  
ACGAACGATCAAGCTGT

### Putative ORFs

20 >-out: 55 to 5997: Frame 1 1981 aa  
 TYLRFGLLYFVAQFISTFGSFLGFHQKQWFHVFVDVLCNEFLATFIVCKIVLFVRHIIVGCNNADCVCACSKS  
 RLKRVPLQTIINGMHKSFYVNANGTCFCNKHNFVCNCDSFGPGNTFINGDIARELGNVVKTAVQPTAPAYVI  
 DVDFVNGFYRLYSGDTFWRYDFDITESKYSCKEVLKNCNVLENFIVYNNSGSNTQIKNACVYFSQLCEPIK  
 VNSELLSTLSVDFNGVLHKAYDVL CNSFFKELTANMSMAECKATLGLTVSDDFVSAVANAHDYDVLLSDLSF  
 NFFISYAKPEDKLSVYDIACCMRAGSKVNVHNVLIKESIPIVVGVKDFNTLSQEGKKVLVTTAKGLTFLLT  
 DNQAITQVPATSIYAKQGAGFKRTYFLWVCLFVVALFIGVSFIDYTTVTFSFHGYDFKYIENGQLKVFEAPL  
 CVRNVDNFNQWHEAKFGVVTNTSDKPIVVGUSERINVVPGVPTNVYLVGKTLVFTLQAAFNGTGVCYDFDGV  
 TSDKCFNSACTRLEGLGGDNVYCNTDILIESKPSILQPNAYKYDVKNYVRFPEILARGFGLRTIRTLATR  
 CRVGECRDHSKGVCFGFDKWVNDGRVDDGYICGDLGLIDLLVNVLSIFSSFSVVAWSGHMLFNFLFAXFITFL  
 FLVTKFKRVFGDLSYGVFTVVCATLNNISYVVTQNLFFMLLYAIIYFVFTRTVRYAWIWHIAVYFLLIPW  
 LLTWFSAFAFLLELPNVFKLKIISTQLFEGDKFIGTFESAAAGTFVLDMRMSYERLINTISPEKLKNYAAASYNK  
 YSGSASEADYRCACYAHLAKAMLDYAKDHNDMLYSPPТИSYNSTLQSGLKKMAQPSGCVERCVVVRVCYGTVLN  
 VWLGDTVTCPRHVIAPSTTVLIDYDHAYSTMRLHNFSVSHNGVFLGVGVTMHGSVLRIVKVSQSNVHTPKHVK  
 LKPGASFNILACYEGIASGVFGVNLRTNTFIKGSFINGACGSPGYNVRNDGTVEFCYLHQIELGSGAHVGSDFT  
 SVYGNFDDQPSLQVESANLMLSDNVVAFLYAAALLNGCRWWLRSTRVNVDGFNEWAMANGTYTIVSSVECYSILA  
 TGVSVQEQLASIQHLHEGFGGNILGYSSLCDDEFLAEEVVKQMYGVNLQSGKVIIFGLKTMFLFSVFFTMFWAELI  
 IYNTNTIWINPVLTPIFC111FLSLVLTMLFLHKFLFLQVFLPTVIALTALYNCVLDYYIVKFLADHFNYNVSI  
 QMDVQGLVNVLVCLFVFLHTWRFSKERFTHWFTYVCSLIAVAYTYFVWFRFFKCTMGVYDFKVSAAEFKYMVANGLHAI  
 LSRLIIFSPESVFSVFGDVKLTLLVYVLIICGYLVCTYWGILYWFNRFFKCTMGVYDFKVSAAEFKYMVANGLHAI  
 YGPFDALWLSFKLLGIGGDRCIKISTVQSKLTDLKCTNVVLLGCLSSMNAANSSEWAYCVDLHNKINLCDDPEI  
 AQGMILLALLAFLFLSKHSDFGLDGLIDSYFDNSSTLQSVASSFVSMPSIAYENAROAYEDAIANGSSSQLIKQ  
 RAMNIAKSEFDHEISVQKKINRMAEQAATQMYKEARSVNRSKVISAMHSLLFGMLRRLDMSVETVNLARDGV  
 VPLSVIPATSASKLTIIVSPDLESYSKIVCDGSVHYAGVWTLNDVKDNDGRPVHVKETRENVETLTWPLLILNCE  
 RRVVKLQNNEIMPGKLQKPKMAEGDGGVLGDDGNALYNTTEGGKTFMYAYISNKADLKFKVWEYEGGCNTIELDSPC  
 RFMVETPNGPQVKLYFVKNLNTLRRGAVLGFIGATIRLQAGKQTELAVNSGLTACAFSVDPATTYLEAVKHGA  
 KPVSNCKMLNSNAGNGQAITTSDVANTNQDSYGGASICLYCRAHVPHPSMDGYCKFKGKCVQVPIGLDPIRFC  
 LENNCNCVCGCWLGHGACDRTTIQSVDILI  
 >-out: 263 to 511: Frame 2 83 aa  
 LVLKVLDLNVYHFKLLLVCINHSMLMLMVVLVSVINITSFVLIVILLGLVILLLMVILQESLVMLLKQLFNPQL  
 LHMLLLIR  
 >-out: 875 to 1054: Frame 2 60 aa  
 LFLMMILFQLLPMHIGMTFCFOICHLIIFLFLMLNLKISCPFMTLVVCPVVLRLLTIMF  
 >-out: 1556 to 1804: Frame 2 83 aa  
 ERLLFLHYRLLLETQVFVMTLMLVPLVISVFLILLVLGWKVWVVTMFIVTTLILLKVLNLIVFYSPLIISMMKL  
 IMYVFQKF  
 >-out: 1808 to 1966: Frame 2 53. aa  
 LEVLAYVLELWLHVIVELVNAVTHIKVFVVLVINGMLMMDVLMVTFTVVMVL  
 >-out: 2600 to 2761: Frame 2 54 aa  
 ITQKIIIMTCYIHLPLATIIPPYNLVLRWHNHLVVLRDVWFASVMVVLCLMEFG  
 >-out: 2798 to 2980: Frame 2 61 aa  
 HHQPLFLLIMIMHIVLCVCIIFQCLIMVSSWELLVLQCMVLCCVLRFHNLMYIHLNMFLKR  
 >-out: 4595 to 4774: Frame 2 60 aa  
 VNTIVILVLMALLILILIIIVAPCRVLLHHLLVCHHILLMKMLDKLMRMLLLMDLLLNLNN  
 >-out: 4790 to 4945: Frame 2 52 aa  
 ISQSLNLIMRYLFRRKLIIEWLNKLLLRCIKKHALLIENLKLVLCTLYFLEC  
 >-out: 5048 to 5200: Frame 2 51 aa  
 LLLVQILNLILRLFVMLVLFIMLELFGHMLKTMVVDLFMLKRLQGRMLKL

~-out: 5753 to 5905: Frame 2 51 aa  
 MLTPIKILMVERLFVCIVGPTFLTLVWMVTVSLRVNVFRFLLVVVWILLGFV

## 5 Alignment

>gi|12175747|ref|NP\_073549.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 gi|30179827|sp|Q05002|R1AB\_CVH22 Replicase polyprotein 1ab (pp1ab) (ORF1ab polyprotein) [Includes:  
 10 Replicase polyprotein 1a (pp1a) (ORF1a)] [Contains: p9;  
 p87; p195 (Papain-like proteinases 1/2)  
 (PL1-PRO/PL2-PRO); Peptide HD2; 3C-like proteinase  
 (3CL-PRO) (3CLp) (M-PRO) (p34); Unknown protein 1; p5;  
 p23; p12; Growth factor-like peptide (GFL) (p16);  
 15 RNA-directed RNA polymerase (RdRp) (Pol) (p100); Helicase  
 (Hel) (p66) (p66-HEL); Unknown protein 2; p41; Unknown  
 protein 3]  
 gi|12082740|gb|AAG48691.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 Length = 6758

20 Score = 2840 bits (7361), Expect = 0.0  
 Identities = 1350/1997 (67%), Positives = 1609/1997 (80%), Gaps = 4/1997 (0%)  
 Frame = +1

25 Query: 10 LDSLFVILVFCNFW\*TYLRFGLLYFVAQFISTFGSFLGFHQKQWFLHFVPPFDVLCNEFLA 189  
 + V+++ F YER LLYFVAQ IST G FLG+ + WFLHF+PFDV+C+E L  
 Sbjct: 2076 MQPFTIVMVLLLIFGDNYLRCFLLYFVAQMISTVGVFLGYKETNWFLHFIPFDVICDELLV 2135

30 Query: 190 TPIVCKIVLFVRRHITVGCNNADCWACSKSARLKRVPQTIINGMHKSFYVNANGGTCFCN 369  
 T IV K++ FVRRH++ GC N DC+A CSKSKSARLKR P+ TI+NG+ +SFYVNANGG+ FC  
 Sbjct: 2136 TVIVVIKVISFVRHVLFGCENPDCIACSKSARLKRFPVNTIVNGVQRSFYVNANGGSKFCK 2195

35 Query: 370 KHNFFCVNCDSFGPGNTFINGDIARELGNVVKTAVQPTAPAYVIIDKVDVFVNGFYRLYSG 549  
 KH FFCV+CDS+G G+TFI +--+RELG+N+ KT VQPT PAYV+IDKV+F NGFYRLYS  
 Sbjct: 2196 KHRFFCVDCDSYGYGSTFITPEVSRELGNIKTNVQPTGPAYVMDKVEFENGFYRLYSC 2255

40 Query: 550 DTFWRYDFDITESKYSCKEVLKNCNVLENFIVYNNSGSNTQIKNACVYFSQLLCEPIKL 729  
 +TFWRY+FDITESKYSCKEVKNCNVL++FIV+NN+G+N+TQ+KNA VYFSQLLC PIKL  
 Sbjct: 2256 ETFWRYNFDITESKYSCKEVKNCNVLDDFIVFNNNGTNVTQVKNASVYFSQLLCRPIKL 2315

45 Query: 730 VNSELLSTLSVDFNGVLHKAYVDVLCSFFKELTANMSMAECKATLGLTVSDDDFVSAVA 909  
 V+SELLSTLSVDFNGVLHKAY+DVL NSF K+L ANMS+AECK LGL++SD +F SA++  
 Sbjct: 2316 VDSELLSTLSVDFNGVLHKAYIDVLRNSFGKDLNANMSLAECKRALGLSISDHEFTSAIS 2375

50 Query: 910 NAHRYDVLLSDLSFNNPFISYAKPEDKLSVYDIACCMRAGSKVVNHNVLIKESIPIVWGV 1089  
 NAHR DVLLSDLSFNNF SYAKPE+KLS YD+ACCMRAG+KVVN NVL K+ PIW  
 Sbjct: 2376 NAHRCDVLLSDLSFNNFVSSYAKPEEKLSAYDLACCMRAGAKVNNANVLKDQTPIVWHA 2435

55 Query: 1090 KDFNTLSQEGKKYLVKTTKAKGLTFLLTFNDNQAITQVPATSIVAKQGAGFK-RTYNFLW 1266  
 KDFN+LS EG+KY+VKT+KAKGLTFLLT N+NQA+TQ+PATSIVAKQGAG + +LW  
 Sbjct: 2436 KDPNSLSAEGRKYIVKTSKAKGLTFLLTINENQAVTQIPATSIVAKQGAGDAGHSLTWLV 2495

60 Query: 1267 YVCLFVVAL-FIGVSFIDYTT--TVTSFHGYDFKYIENGQLKVFEAPLHCVRNVFDNFNQ 1437  
 +C V + F F+ Y V+SF GYDFKYIENGQLK FEAPL CVRNVF+NF  
 Sbjct: 2496 LLCGLVCLIQFYLCFFMPYFMYDIVSSFEGYDFKYIENGQLKNEAPLKCVRNVFENFED 2555

65 Query: 1438 WHEAKFGVVTINSKCPIVVGVSERINVVPGVPTNVYLVGKTLVFTLQAAFGNTGVCYDF 1617  
 WH AKFG N CPIVVGSE +N V G+P+NVYLVGKTL+FTLQAAFGN GVCYD  
 Sbjct: 2556 WHYAKFGFTPLNKQSCPIVVGVSIEVNTVAGIPSNSVYLVGKTLIFTLQAAFGNAGVCYDI 2615

70 Query: 1618 DGVTTSKDKCIFNSACTRLEGLGGDNVYCYNTDLIEGSKPYSILQPNAAYKYDVKNYVRFP 1797  
 GVTT +KCF SACTRLEGLGG+NVYCYNT L+EGS PYS +Q NAYYKYD N+++ P  
 Sbjct: 2616 FGVTTPKEKCIITSACTRLEGLGGNNVYCYNTALMEGSLPYSSIQANAYYKYDNGNFIKLP 2675

75 Query: 1798 EILARGFGLRTIRTLATRYCRVGECRDSHKGVCFGFDKWWVNDGRVDDGYICGDDGLIDXX 1977  
 E++A+GFG RT+RT+AT+YCRVGE +S+ GVCFCFDKW+VNDGRV +GY+CG GL +  
 Sbjct: 2676 EVIAQGFGFRTVRTIATKCYCRVGEVESNAGVCFGFDKWVNDGRVANGYVCGTGLWNLV 2735

80 Query: 1978 XXXXXXXXXXXXXXXAMSGHMLFNFLFAXFITFLCFLVTKFKRVFGDLSYGVFTVVVCATLI 2157  
 AMSG +L N F F CFLVTKF+R+FGDLS GV TVV A L+  
 Sbjct: 2736 FNILSMFSSSFSAAMSGQILLNCALGAFIAFCCFLVTKFRRMFGDLSVGVCVVAVLL 2795

85 Query: 2158 NNISYVVTQNLFFMLLYAILYFVFTRTVRYAWIWHIAVIVAYFLLIPWWLLTWFSFAAFL 2337  
 NN+SY+VTQNL M+ YAILYF TR++RYAWIW AY++AY PWWL W+ A

Sbjct: 2796 NNVSYIVTQNLVTMIAVAILYFFATRSLRYAWIWCAAYLIAYISFAPWWLCAWYFLAMLT 2855  
 Query: 2338 ELLPNVFKLKJSTQLFEGDKFIGTFESAAAGTFVLDMRSYERLINTISPEKLXXXXXXX 2517  
 5 Sbjct: 2856 GLLPSLLKLKVSTNLFEGDKFVGTESAAAGTFVIDMRSYEKLANISISPEKLKSYAASYN 2915  
 Query: 2518 XXXXXXXXXXXXEAODYRCACAYAHLAKAMLDYAKDHNDMLYSPPSTISYNSTLQSGLKKMAQPS. 2697  
 EADYRCACAYA+LAKAMLD+++DHND+LY+PPT+SY STLQ+GL+KMAQPS  
 10 Sbjct: 2916 RYKYYSGNANEADYRCACAYAHLAKAMLDFSRDHNDILYTPPTVSYGSTLQAGLRKMAQPS 2975  
 Query: 2698 GCVERCVVRVCYGSTVNLNGWLGDTVTCPRHVIAPSTTVLIDYDHAYSTMRLHNFSVSHN 2877  
 G VE+CVVRVCYGV+TVLNG+WLGD V CPRHVIA +TT IDYDH YS MRLHNFS+  
 Sbjct: 2976 GFVEKCVVRVCYGNLTVNLGLWLGDIVYCPRHVIASNTTSAIDYDHEYSIMRLHNFSIISG 3035  
 15 Query: 2878 GVFLGVVGVVTMHGSVLRIKVSQSNVTPKHFVTLKPGASFNLACYEGIASGVFGVNLR 3057  
 FLGVVG TMHG L+IKVSQ+N+HTP+H F+TLK G FNILACY+G A GVFGVN+R  
 Sbjct: 3036 TAFLGVVVGATMHGVTLKIKVSQTNMHTPRHSFRTLKSGEFGNIFLACYDCAQGVFGVNM 3095  
 20 Query: 3058 TNFTIKGSFINGACGSPGYNVRNDGTVEFCYLHQIELGSGAHVGSDFTGSVYGNFDDQPS 3237  
 TN+TI+GSFINGACGSPGYN++N G VEF Y+HQIELGSG+HVGS F G +YG F+DQP+  
 Sbjct: 3096 TNWТИRGSFINGACGSPGYNLKN-GEVEFVYMHQIELGSGSHVGSSFDGVMYGGFEDQPN 3154  
 Query: 3238 LOVESANMLSNDNVVAFLYAAALLNGCRWWLRSTRVNVDGFNEWAMANGYTIVSSVECYSI 3417  
 25 Sbjct: 3155 LOVESAN ML+ NVVAFLYAA+LNGC WWL+ ++ V+ +NEWA ANG+T ++ + +SI  
 LOVESANQMLTVNVVAFLYAAAILNGCTWWLKGELFVHEHNEWAQANGFTAMNGEDAFSI 3214  
 Query: 3418 LAAKTGVSVQQLLASIQHLHEGFGGKNIILGYSSLCDDEFTLAEVVKQOMGVNLQSGKVIFG 3597  
 LAAKTGV VE+LL +IQ L+ GFGGK ILGYSSL DEF++ EVVKQOM+GVNLQSGK  
 30 Sbjct: 3215 LAAKTGVCVERLLHAIQVLNNNGFGGKQILGYSSLNDEFSINEVVVKQOMGVNLQSGKTTSM 3274  
 Query: 3598 LKTMFLFVFFTMFWAELFIYTNTIWINPVIXXXXXXXXXXXXXXXXXXXXXXXHHKFLLQVF 3777  
 K++ LF+ FF MFWAELF+YT TIW+NP KHK LFLQVF  
 Sbjct: 3275 FKSISLFAFPFVFMFWAELFVYTTTIWVNPGLTPFMILLVALSLCLTFVVKHKVLFQVF 3334  
 35 Query: 3778 LLPTVIATALYNCVLDYYIVKFLADHFNYNVSVLQMDVQGXXXXXXXXXXXXXXHTWRF SK 3957  
 LLPP++I A+ NC DY++ K LA+ F+YNSV+QMD+QG HTWRF+K  
 Sbjct: 3335 LLPSIIVAAIQNCANDYHVTKVLAEKFDYNVSVVMQMDIQGFVNIFICLFLVALLHTWRF A 3394  
 Query: 3958 ERFTHWFTYVCSLIAVAYTYFSGDFLSLLVMFLCAISSDWYIGAIVFRLSRLIIFFSPE 4137  
 40 ER THW TY+ SLIAV YT YS D++SLLVM LCAIS++WYIGAI+FR+ R + F P  
 Sbjct: 3395 ERCTHWCTYLFSLIAYLTYALSYDVVSLLVMLLCAISNEWYIGAIIFRICRFGVAFPLV 3454  
 Query: 4138 SVFSVFGDVKITLVVYILICGYLVCTYWGILYWPNRFFKCTMGVYDFKVSAAEFKYMVANG 4317  
 S F VK L+ Y++ G++ C Y+G+LYW NRF KCT+GVYDF VS AEFKYMVANG  
 45 Sbjct: 3455 EYVSYFDGVKTVLLFYMLLGFVSCMYGLLYWINRFCKCTLGVYDFCVPSPAEFKYMVANG 3514  
 Query: 4318 LHAPYGPFDALWLSFKLLGIGGDRCIKISTVQSKLTLKCTNVVLLGCLSMMNTIAANSSE 4497  
 L+AP GPFDAL+LSFKL+GIGG R IK+STVQSKLTLKCTNVVL+G LS+MNIA+NS E  
 50 Sbjct: 3515 LNAPNGPFDALFLSFKLMGIGGPRTIKVSTVQSKLTLKCTNVVLMGILSNMNTIASNSKE 3574  
 Query: 4498 WAYCVDLHNKINLCCDPKAQGMILLALLAFLSKHSDFGLDGLIDSYFDNSSTLQSVASS 4677  
 WAYCV++HNKINLCCDP AEQ +LLALLAFLSKHSDFGL L+DSYF+N S LQSVASS  
 Sbjct: 3575 WAYCVMHNKINLCCDPETAQELLALLAFLSKHSDFGLDVLDSYFENDSILQSVASS 3634  
 55 Query: 4678 FVSMPSYIAYENARQAYEADAIANGSSSQLIKQLKRAMNIAKSEFDHEISVQKKINRMAEQ 4857  
 FV MPS++AYE ARQ YE+Ä+ANGSS Q+IKQLK+AMN+AK+EFD E SVQKKINRMAEQ  
 Sbjct: 3635 FVGMPSPFVAYETARQAYENAVANGSSPQIICKQLKKAMNVAKAEFDRESSVQKKINRMAEQ 3694  
 60 Query: 4858 AATOMYKEARSVRNSKSVIISAMHSLLFGMLRRLDMSSVETVNLARDGVVPLSVIPATSA 5037  
 AA MYKEAR+VNRNSKSV+SAMHSLLFGMLRRLDMSSV+T+LN+AR+GVVPLSVIPATSA  
 Sbjct: 3695 AAAAMYKEARAVNRNSKSVSAMHSLLFGMLRRLDMSSVDTILNMARNGVWPLSVIPATSA 3754  
 Query: 5038 SKLTIVSPDLESYSKIVCDGSVHYAGVVWTLNDVKDNDGRPVHVKEITRENVETLTWPLI 5217  
 65 Sbjct: 3755 ARLVVVVVPDHDSFVKMMVDFVHYAGVVWTLQEVKDNDGKVNHLKDVTKENQEILVWPLI 3814  
 Query: 5218 LNCERVVKLQNNEIMPGKLKQKPMKAEGDGGVLGDGNALYNTTEGGKTFMÝAYISNKA DLK 5397  
 L CERVVKLQNNEIMPGK+K K K EGDGG+ +CNALYN EGG+ FMYAY++ K +K  
 70 Sbjct: 3815 LTCERVVKLQNNEIMPGKMKVKATKGEGDGGITSEGNALYNNEGGRAFMYAYVTTKPGMK 3874  
 Query: 5398 FVKWBEYEGGCNTIELDSPCRFMVETPNPGPQVKYL FVKNLNTLRRGAVLGFIGATIRLQA 5577  
 +VKWE++ G T+EL+ PCRF+++TP GPQ+KYLYFVKNLN LRRGAVLGFIGATIRLQA  
 Sbjct: 3875 YVKWEHDSGVVTVELEPPCRFVIDTPGPQIKYLYFVKNLNNLRRGAVLGFIGATVRLQA 3934  
 75 Query: 5578 GKQTELAVNSGLLTACAFSVDPATTYLEAVKHGAKPVSNICMILSNGAGNGQAITTSVDA 5757  
 GKQTE NS LLT C+F+VDPA YL+AVK GAKPV NC+KML+NG+G+GQAIT ++D+

Sbjct: 3935 GKQTEPVSNHLLTHCSFAVDPAAAYLDAVKQGAKPVGNCVKMLTNGSGSGQAITCTIDS 3994

Query: 5758 NTTNQDSYGGASICLYCRAHVPHPSMDGYCKFKGKCVQVPIGCLDPIRFCLENNVCVCGC 5937  
NTI QD+YGGAS+C+YCRAHV HP+MDG+C++KGK VQVPIG DPIRFCLEN VC VCGC

5 Sbjct: 3995 NTTQDTYGGASVCIYCRAHVAHPTMDGFCQYKGKWWQVPIGTDPIRFCLIENTVCKVCGC 4054

Query: 5938 WLGHGCACDRRTIQSVD 5988  
WL HGC CDRT IQS D

Sbjct: 4055 WLNHGCTCDRTAIQSFD 4071

10

**4. Sequence D**

5325 nucleotides; Replicase

TAGCTTGATTCTGTCGAGCAAGGGTTCTAGTGCAGCTCGACTAGAACCTGTAATGGCACGGACATCGATAAGTG  
 15 TGTTCTGCTTTGACATTATAATAAATGTTTCATTCTTGGGTAAGTGTGAAAGATGAACGTGTTGTCGTT  
 TAAAAATGCTGATCTTAAGGATGGTTATTTGTTATAAAAGAGGTGACTAAGTCGGTTATGGAACACGAGCAATC  
 CATGATAAACCTACTTAACCTTCTGGGCTTGGCTGAGCATGATTCTTACTTGGAAAGATGGCAGAGTCAT  
 TTATGGTATGTTAGTAGACATAATCTTACTAAATATACTATGATGGACTTGGTTATGCTATGCGTAACCTTGA  
 20 TGAACAAAATGTTGATGTTCTAAAAGAAGTATTAGTTAACCTGGTGTGACAATTCTTATTGATAGTAA  
 GGGTGGTATGACCCAGTTGAAAATGAAAGATATACTAGAGTTATGCATCTTGGAAAATTGTTAGCTAGAGC  
 TATGCTTAAATGCGTTGCTCTATGTGATGCGATGGTTGCTAAAGGTGTTGTTGGTCTTAACTAGATAACCA  
 AGATCTTAAATGGTAACTTTATGATTTGGTATTTGGTGTAGCTTACCTAATATGGGTGTTCCCTGTTGAC  
 ATCATATTATTCTTATATGATGCCATTATGGTTAACTAATTGTTAGCTAGTGTAGTGTGTTGGTCAAGAGTGA  
 25 TATTGGTAGTGATTAAAACCTTTGATTTGCTTAAGTATGTTACTGACCTTAACTGAAACATAAAAGAAATTATTCAA  
 TAAGTACTTTAACGATTGGAGTTTGATTTGATTATCATCTTAAATTGTTAGCTAGTGTATGATGATATGTTGTTATACA  
 TTGCTAATTAAACACTATTGCTTACACTATACAGGTTACTGCTTGGTCTACATGCTAAAGTTTT  
 TATAGATGGTGTCCACTTGTACAACACTGCTGGTTATCATTTAACAAATTAGGTTGGGAAATAAGATGT  
 TAACACACACTCAGTTAGGTTGACAATCACTGAACCTTGCCTTACTGACCCCTCCTGATAATAGCTC  
 TTCTCCAGCAGCTGTTGATCAACGACTATTGTTTCTGTTGAGCATTGAGTACTGGTTGACAATCAAGT  
 30 TGTTAAGCCAGGTCAATTAAATGAAAGAGTTTATAACTTCTTGTAAAGAGGTTCTTGATGAAAGGTTCTGA  
 ACTTACATTAAAACATTCTTCTTCGACAGAAATGGTGTGCTGTTAAAGATTTGACTTTACCGTTATAA  
 TAAGCCTACCATTAAAGGATATTGTCAGCTAGAGTACATATAAGATAGTCTCTCGTTATTGACATTATGTA  
 AGGTGGCTGTATTAAAGGCATGTGAAGTGTGTTAAGCAAAATCTTAAAGAGTGTGTTGGCATTAAATAAGTT  
 TGGTAAAGCTAGTTGTTACGATCTATATCTTAAAGAAACAGGATGCTTGTGTTGACAAGCGTAA  
 35 TGTCCCTCCTACTATGACACAGCTGAACTTAAAGTGTGCTTAAAGAACAGTGTGCTAGAAGCTGTTGG  
 TGTTCTCTGTTCCACAATGACCAAGACAATACCATCAAAACATCTTAAACCCATTGTTAACACCGCAA  
 TGCCACTGTTGTTATTGGTACTACCAAAATTGTTGGGGTTGGAATAATATGTTGCGTACTTTAACATTGATGGTGT  
 TGAAAACCTATGCTATGGGTTGGGATTATCCAAATGTGATAAGGCTTGCCTAACATGATAACGTATGATTTC  
 AGCCATGGTGTGGGTTCTAAGCATGTTAATTGTTGACTGTAACAGATAGGTTTATAGGCTTGGTACGACTCTGGTGA  
 40 GGCACAAGTTAACAGAAAGTTGTTATTCTAATGGTGTGTTTTTAAACATTTCAGCCGGTGTAGCTAACATTGCTATAGGTT  
 CGCTAGTACAGCTTATGCTAATTCTTATGAGTACTTCTAAATGTTGGGTTGAAAGAAAGATTAAACTAAGGACC  
 TGTCCTCATGAGTTGTTCCAGCATACTATGCAAATAGTTGATAAAAGATGGTACTCTTATGCTTACCCAGATCC  
 TAGTAGGATCTTGTCAAGCTGGTTTTGTTGATGATGTTGTAAGCAGATGCTGTTGGTGTAGATAATTGCTATAGGTT  
 AACTAGTGTGAAAGAGTCATTGATGATTATTGTTATCTTAAAGGTTTACATTGCTATGATGATTCTCTC  
 TGATGACGGTGTGTGTTATAACAAGGATTATGCTGAGTTAGGTTATAGCAGACATTAGTGTGTTAAAGC  
 45 CACTTGTATTACCAAGATAATGTTTATGAGTACTTCTAAATGTTGGGTTGAAAGAAAGATTAAACTAAGGACC  
 ACATGAGTTTGTGTTCCAGCATACTATGCAAATAGTTGATAAAAGATGGTACTCTTATGCTTACCCAGATCC  
 TAGTAGGATCTTGTCAAGCTGGTTTTGTTGATGATGTTGTAAGCAGATGCTGTTGGTGTAGATAATTGCTATAGGTT  
 TGTGTCATTAGCTATTGATGCTACCCCTCTTCAAAACACCCATTCTGTAAGGTTTTCTGTTACACTTCTGTA  
 ACTGATGGGTTAACGATCTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTT  
 50 TAATCAAGAAAGATAAGTTTGGTGTGAAGGATTTTATGCTAGTATGTTGAAAGCTTCTGTTGCGTAAGCTATGTTGAC  
 TGGCTTATGTTGTTGTTGTTCAAAACTGTTCTCGTTGTTGCGTAAGCTATGTTGAC  
 TAAATGTCATATGATCATGTTGGTACCGACCACAGTTTATTTGGCTATAACACCGTATGTTGTAATG  
 ATCAGGTTGTTGTTAGTGTGTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTTTAAAGGTT  
 55 TGTTGAAGTTTAATAGGCTTGCACGCTGATGGACTGTTAGGGACTATAAACCTGCTAATGTTGAA  
 AGATACACTTAGACTCTTGCCTGGTCAAAAGGTTTAAAGGTTTAAAGGTTAAGGTTAAGGTTAAGGTTAAGGTT  
 AACTCTTAAAGGGTTGTTGGACCTAAAGAATTGCTTCTTAGTTGGGAAAGGTGTTAAAGGTTAAGGTTAAGGTT  
 TCGTAATTCTGTTTCACTGTTTCAAAATAAGTAAAGGACTCAAAATTCCAAATAGGTGAGTTCATCTTGA  
 60 GGTGTAATGTTGTTGTTGACTGTTACGTTACGTTAAAGTCTGTAACCAACTAAGTTAGTCTCTGGTATGATTGTT  
 CTTAACATCTCACAATGTTCAACCTTACGTCACCAACTATTGCAAACCAAGAGAAGTATTCTGCTTAAACAAAA  
 ATTGCAACCTGCTTTAATGTCAGTGTGATGCAATATGCTAATTGTTCCATTACCAACTTATTGGTAAACAAAA  
 GATAACTACAATACAGGGTCTCTGGTAGGGTAAGTCACATTGTTCCATTGGACTTGGATTGACTATCCAGG  
 TGGCGTATTGTTGTTGCTTGTGCCCAGCTGCTGTTGATTCTTATGTCGAAAAGCTATGACTGTTTATAG  
 CATTGATAAGTGTACTAGGATTATACCTGCAAGAGCTCGGGTTGAGTGTAAATGCTGATATTGTTGTTAGTGAAGT  
 65 TAGTGCACAAATACATATTAGCACTGTTAACGCACTTACCTGAGTGTAAATGCTGATATTGTTGTTAGTGAAGT  
 TTCAATGTCACAAATTATGACCTTCTGTTATTAAATCAGCTTATCATATAAACATATTGTTTATGTTGTTAGTGA  
 TCCACAAACAACCTCCTGCAACCTAGAGTAATGATTACTAAAGGTTGTTATGGAGCCTGTTGATTATAACGTTGTTAC  
 TCAACGTTATGTCATAGGCCCCATGTTTCTTCAAAATGTTAGATGTCCTGCTGAAATAGTTAATAC

AGTTTCTGAACTGTTATGAGAACAAAGTTGCCCCGTTAACCTGCTAGTAAACAGTGTAAAATCTTTT  
TAAGGGTAATGTACAGGTTGACAATGGCTCTAGTATTAACAGAAAGCAGCTGAAATAGTTAAGCTGTTTTAGT  
AAAAAATCCAAGGTTGAGTAAGGCTGTTTATTTCTCTTATAATAGTCAGAATTATGTTGCTAGTAGATTTT  
AGGACTTCAAATTCAAACGTGATTCTCTCAAGGTAGTGAGTATGATTATGTAATCTATGCACAAACTCTG  
5 CACTGCACATGCTTGCATGAAACGTTTAATGTTGCTATAACACGTGCTAAGAAGGGTATATTGTTGTAAT  
GTGTGATAAAACTTTGTTGATTCACTTAAGTTTTGAGATTAACATGCAGATTACACTCTAGCCAGGTTG  
TGGCTGTTAAAAAATTGTACACGCACCTCTTAATTACCAACCTCATGCACACACTTCTTGTGCTGTTG  
AGATCAGTTAACACTACAGGTGATTTAGCTGTTCAAATAGTTCAAATAATGTTGACTTATGAACATGTTA  
10 ATCATTTATGGGTTTGTAGGTTGATATTAGTATTCTGGTAGTCATAGTTGTTTGTACACGTGACTTTGCTA  
TCGTAAATGTGCGGGTGGGTGGATGTTGAAAGTGCATGTTGTCAGGCGATAACATAGGTACTAATG  
TCCTTACAGGTTGGTTTCAATGGTGTAAATTGTTGTCAGGACTGAAGGTTGTTGCTACCAATTG  
TGATGTTATTAACCTGTTGCAAAATCTCCACCAAGGTGACAATTAGACACCTTGTCTTACGTA  
AGGACAACCTGGTTAATTGTTGCTAGACGCATTGTGCAAATGATATCTGATTATTGTCACATTGCTGACAT  
15 TCTTGTCTTGTGTTGGGCAAGGTAGTTGGAATTAACTACAATGCGTTACTTTGTAAAAATAGGGCAATT  
ATATTGTTATGTGGTAATTCTGCCACTTGTATAATTCACTGTTGAGTACACAGTGGGGTTATGTTGGTTCTTGAGGCCAGA  
GGGTGTTGATTATGTTACAATCCGTATGCTTTGATACACAGTGGGGTTATGTTGGTTCTTGAGGCCAGA

### Hypothesized ORFs

>-out: -1 to 5320: Frame 2 1774 aa  
SLIRRARGSSAARLEPCNGTDIDKCVRAFDIYNKNVSFLGKCLKMNCVRFKNADLKDGYFVIKRCTKSVMEHEQS  
MYNLLNFSGALAEHDFFTWDGRVIYGNVSRHNLTKYTMMMDLVYAMRNFDEQNCVLKEVLVTGCCDNSYFDSI  
GWYDPVENEEDIHRVYASLGKIVARAMILKVALCDAMVAKGVVGVLTLNDNQDNGNFYDFGDFVVSPLNMGVPCCT  
25 SYYSYMPIMGLTNCLASECFVKSDFKTFDLYKDFTEHKENLFNKYFKHWSFDYHPNCSDCYDDMCVIF  
CANFNTLFATTIPTGTAFCPLCRKVFDGVPVTTAGYHFKQQLGLVWNKDVNTHSRLTITELLQFVTDPSLIIAS  
SPALVDQRTICFSVAALSTGLTNQVVKPGHNEEFYFNFLRLRGFFDEGSELTLKHHFFAONGDAAVKDFDFYRYN  
KPTILDICQARVTVYKIVSRYFDIYEGGCIKACEVVVTNLNSAGWLNKGKASLYYESISYEEQDALFALTKRN  
VLPTMTQLNLKYAISGKERARTVGGVSLLSTMTRQYHQHKLKSIVNTRATVVIQVTTKFYGGWNNMLRTLIDGV  
ENPMLMGWDYPKCDRALPNMIRMIISAMVLGSKHVNCCVTDRYFRLGNELAQVLTEVVYNSGGFYFKPGGTTSGI  
30 ASTAYANSIFNIFQAVSSNINRLLSVPSDSCNNVNRDLQRRLYDNCYRLTSVEESFIDDYYGYLKRHFSSMILS  
DDGVVCYNDYAEGLYIADISAFKATLYYQNNVFMSTSCKWVEEDLTKGPHFCSQHTMQIVDKDGTYYLPYPD  
SRILSAGVFVDDVVKTDAVVLERYVSLAIDAYPLSKHPNSEYRKVFYVLLDWFVHLNKNLNEGVLESFSVTLLI  
NQEDKFWCEDFYASMYENSTILQAAGLCCVCGSQVLRCGDCLRKPMLCTKACAYDHVFGTDHKFILAIPYVCNA  
SGCGVSDVKKLYLGGLNYYCTNMHKPQLSPLCASAGNIFGLYKNSATGSLDVEVFMRLATSDWTDVRDYKLANDV  
35 DTLRLFAAETIKAKEESVKSSYAFATLKVNVPKKELLSWESGKVKPPLNRNSVFTCFQISKDSKFQIGEFIFER  
VEYGSDTVTYKSTVTKLVPGMIFVLTSHNVGPKLRAPICTANEKYSSYIYKLHPAFNVSDAYANLVPYYQLIGKQK  
ITTIQGPPGSGKSHCSIGLGLYYPGARIYFVACAHAAVDSLCAKAMTVYSIDKCTRIIPARARVECYSGFKPNNT  
SAQYIFSTVNALPECNAIVVDEVSMTNYDLSVINQRLSYKHIVYVGDPQQLPAPRVMITKGVMEPVDVNVVI  
40 QRMCAIGPDVFLHKCYRCPAEIVNTVSELVYENKFPVVKPASKQCFKIFFKGNVQVDNGSSINRKQLEIVKLFV  
KNPSWSKAVFISPNSQNYVASRFLGLQIQTVDSSQGSEYDVYIYAQTSDTAHACNVNRFNVAITRAKKGIFCVM  
CDKTLFDSLKFFEIKHADLHSQVCGLFKNCRTPLNLPPTAHTFLSLSDQFKTTGDLAVQIGSNNVCTYEHVI  
SFMGFRFDISIPGSNSLFLCTRFAIRNVRGWLGMVESAHVCGDNIGTNVPLQVGFNSNGVNFVQTEGCVSTNFG  
DVIKPVCAKSPPGEQFRHLVPFLRKQGPWLIVRRRIVQMSIDYLSNLSDILVFVWLWAGSLELTMRYFVKIGPIK  
YCYCGNSATCYSVSNEYCCFKHALGCDYVYNPYAFDIQQWGYVGSLSQ  
45 >-out: 189 to 341: Frame 3 51 aa  
RGVLSRLWNTSNPCITYLTFLVLWLSMISLLGKMAESFMVMLVDIILNL  
>-out: 726 to 977: Frame 3 84 aa  
LVSVLSRVIFLVVILKLLICLSMISLNIKKIYSISTLSIGVLIILIVVTVMMICVLYIVLILIHYPQLYQVLLLV  
HYVVFKL  
50 >-out: 2661 to 2903: Frame 3 81 aa  
MRVFLNLFLHHFLIIKKISFGVKIFMLVCMKILQYCKLLAYVLFVVHKLFFVVVIVCVSLLCCALNVHMIMY  
VPTTSFLFWL  
>-out: 3075 to 3296: Frame 3 74 aa  
MLKFLIGLQLRIGLMLGTINLLMMLKIHLDLSRLKLLKLRVLSLLMLLQLLKRLLDLKNCFLVVGKVVK  
LNHL  
55 >-out: 3741 to 3890: Frame 3 50 aa  
LFIALISVLGLYQELGLSVIVALNQITLVHNTYLALLTHYLSVMLLLL  
>-out: 4500 to 4676: Frame 3 59 aa  
CVIKLCLIHLSFLRLNMQIYTLARFVACLKIVHALLIYHQLMHTLSCRCQISLRLQVI  
60 >-out: 4692 to 4862: Frame 3 57 aa  
VQIMFVLMNMLYHLLWVGLLIVFLVVIVCFVHVTLLFVMCVVGWVWMLKVLMFVAIT  
>-out: 4866 to 5039: Frame 3 58 aa  
VLMFLYRLVLFQMVLILLCKLKVCLPILVMLLNLFVQNLHQVNNLDTLFLFYVKDNLG  
>-out: 5166 to 5315: Frame 3 50 aa  
GQLNIVIVVILPLVIIQLVMNTIVVLMNHWWVIMFTIRMLLIYNSGVMLVP

## Alignment

>gi|12175747|ref|NP\_078549.1| replicase polyprotein 1ab [Human coronavirus 229E]  
>gi|30179827|sp|Q05002|R1AB\_CVH22|Replicase polyprotein 1ab (pp1ab) (ORF1ab polyprotein) [Includes: R1AB\_CVH22]

5 Replicase polyprotein 1a (pp1a) (ORF1a)] [Contains: p9;  
p87; p195 (Papain-like proteinases 1/2)  
(PL1-PRO/PL2-PRO); Peptide HD2; 3C-like proteinase  
(3CL-PRO) (3CLp) (M-PRO) (p34); Unknown protein 1; p5;  
p23; p12; Growth factor-like peptide (GFL) (p16);  
10 RNA-directed RNA polymerase (RdRp) (Pol) (p100); Helicase  
(Hel) (p66) (p66-HEL); Unknown protein 2; p41; Unknown  
protein 3]

protein 31  
gi|12082740|gb|AAG48591.1| replicase polyprotein 1ab [Human coronavirus 229E]  
Length = 6758

15 Score = 3137 bits (8134), Expect = 0.0  
Identities = 1465/1773 (82%), Positives = 1633/1773 (92%)  
Frame = +2

20	Query: 2	SLIRRARGSSAARLEPCNGTIDKCVRAFDIYNKNVSFLGKCLMNCVRFKNADLKDGYF 181
	Sbjct: 4073	S + R RGSSAARLEPCNGTID CVRAFD+Y NK+ SF+GK LK NCVRFKN D D ++ SYLNRVRGSSAARLEPCNGTIDYCVRAFDVYNKDASFIGKNLKSNCVRFKNVDKDDAFY 4132
25	Query: 182	VIKRCKTSVMEHEQSMYNNLNFSGALAEHDFFTWDKGVRVYGNVSRHNLTKYTMMDLVYA 361
	Sbjct: 4133	++KRC KSVM+HEQSMYNNL A+A+HDFFTW +GR IYGNVSR +LTKYTMMDL +A IVKRCIKSVMDEQSMYNNLKGCAVAKHDFFTWHEGRTIYGNVSRQDLTKYTMMMDLCFA 4192
30	Query: 362	MRNFDQEQNCDVLKEVLVLTGCCDNSYFDISKWYDPVENEEDIHRVYASLGKIVARAMLKCV 541
	Sbjct: 4193	+RNFDE++C+V KE+LVLTGCC YF+ K W+DP+ENEEDIHRVY+LGK+VA AMLKCV LRNFDEKDCEVFKIEILVLTGCCSTDYFEMKNWFDPINEEDIHRVYAA LGKVVANAMLKCV 4252
35	Query: 542	ALCDAMVAKGVGVGVLTLDNQDLNGNFYDFGDFVVSPLPNMGVPCCSTSYYSYMPIMGLTNC 721
	Sbjct: 4253	A CD MV KGVVGVVLTDNQDLNGNFYDFGDFV+ P MG+P CTSYYSYMP+MG+TNC AFCDEMVLKGVVGVLTLDNQDLNGNFYDFGDFVLCPPGMGIPYCTSYYSYMPVGMGTMNC 4312
40	Query: 722	LASECFVKSDIFGSDFKTDFLLKYDFTEHKENLFLNKYFKHWSFDYHPNCSDCYDDMCVIH 901
	Sbjct: 4313	LASECF+KSDIFG DFKTFDLLKYDFTEHKE LFNKFK+W DYHP+C DC+D+MC++H LASECFMKSDFQDFKTFDLLKYDFTEHKEVLFNKFYKFWGQDHYHPDCVDCHDEMCLIH 4372
45	Query: 902	CANFNTLFATTIPIGTAFGPLCRKVFIGDGVPLVTTAGYHFKQLGLVWNKDVNTHSVRLTIT 1081
	Sbjct: 4373	C+NFTLFATTIPI TAFGPLCRKVFIGDGV+V TAGYHFKQLGLVWNKDVNTHS RLTIT CSNFTLFATTIPINTAFGPLCRKVFIGDGVVVATAGYHFKQLGLVWNKDVNTHSRTRLTIT 4432
50	Query: 1082	ELLQFVTDPSLIIASSPALVDQRTICFSVAALSTGLTNQVVKPGHFNEEFYNFLRLRGFF 1261
	Sbjct: 4433	ELLQFVTDPSLIIASSPALVD+RT+CFSVAALSTGLT+Q VKPGHFN+EFY+FLR +GFF ELLQFVTDPTLIVASSPALVDKRTVCFSVAALSTGLTSQTVKPGHFNKEFYDFLRSQGFF 4492
55	Query: 1262	DEGSELTLKHFQQAQNGDAAVKDFDFYRYNKPFIIDICQARVTVYKIVSRYFIDYEGGCIC 1441
	Sbjct: 4493	DEGSELTLKHFQQAQNGDAAVKDFDFYRYNKPFIIDICQARVTVYKIVSRYFIDYEGGCIC DEGSELTLKHFQQAQNGDAAVKDFDFYRYNKPFIIDICQARVTVYKIVSRYFIDYEGGCIC 4552
60	Query: 1442	ACEVVVTNLNKSAGWPLNKGKASLYYESISYEEQDALFALTKRNLVPTMTQLNLKYAIS 1621
	Sbjct: 4553	+ EVVVTNLNKSAGWPLNKGKASLYYESISYEEQDALF+L+LTKRN+LPTMTQLNLKYAIS SREVVVTNLNKSAGWPLNKGKAGLYYESISYEEQDAIFSLTKRNILPTMTQLNLKYAIS 4612
65	Query: 1622	GKERARTVGGVSLLSTMTTRQYHQKHLKSIVNTRNATVVIGTTKFYGGWNMLRTLIDGV 1801
	Sbjct: 4613	GKERARTVGGVSLL+TMTTRQ+HQK LKSIV TRNATVVIGTTKFYGGW+NML+ L+ V GKERARTVGGVSLLATMTTRQFHQKCLKSTIVATRNATVVIGTTKFYGGWDNMLKNLMAADV 4672
70	Query: 1802	ENPMLMGWDYPKCDRALPMIRMISAMVLGSKHVNCCTVTDRFYRLGNELAQVLTEVVYS 1981
	Sbjct: 4673	++P LMGWDYPKCDRA+P+MIRM+SAM+LGSKHV CCT +D+FYRL NELAQVLTEVVYS DDPKLMGWDYPKCDRAMPMSIRMLSAMILGSKHVTCCTASDKFYRLSNEAQVLTEVVYS 4732
75	Query: 1982	NGGFYFKPGGTTSGDASTAYANSIFNIFQAVSSNINRLLSVPSDSCNNVNVRDLQRRLYD 2161
	Sbjct: 4733	NGGFYFKPGGTTSGDASTAYANS+FNIFQAVSSNIN +LSV S +CNN NV+ LQR+LYD NGGFYFKPGGTTSGDASTAYANSFVNIFQAVSSNINCVLSVNSSNCNNFNVKKLQRQLYD 4793
80	Query: 2162	NCYRLTSVEESFIDDDYYGYLKRHSMMILSDDGVVCYNKDYAELGYIADISAFKATLYYQ 2341
	Sbjct: 4793	NCYR ++V+ESF+DD+YGYL+KHFSSMILSDD VVCYNK YA LGYIADISAFKATLYYQ NCYRNSNVDESVDFFYGYLQKHFSSMILSDDSVVCYNKTYAGLGYIADISAFKATLYYQ 4851
85	Query: 2342	NNVFMSTSKCWEEDLTKGPHEFCSQHTMQIVDKDGTYYLPPDPSRILSAGVFVDDVVK 252
	Sbjct: 4853	N VFMST+KCW EEDL+ GPHEFCSQHTMQIVD+G YYLPYPPDPSRI+SAGVFVDD+ K NGVFMSTAKCWTEEDLSIGPHEFCSQHTMQIVDENGKYYLPPDPSRIISAGVFVDDITK 4911

Query: 2522	TDAVILLERYVSLAIDAYPLSKHPNSEYRKVFYVLLDWVKHLNKNLNEGVLLESFSVTLLD	2701
Sbjct: 4913	TDAVILLERYVSLAIDAYPLSKHP EYRKVFY LLDWKHLNK LNEGVLLESFSVTLLD	4972
5	Query: 2702 NQEDKFWCEDFYASMYENSTILQAAAGLCVVCGSQTVLRCGDLRKPMCLTKCAYDHVFGT E KFW E FYASMYE ST+LQAAGLCVVCGSQTVLRCGDLR+PMLCTKCAYDHVFGT	2881
Sbjct: 4973	EHESKFWDESFYASMYEKSTVLQAAAGLCVVCGSQTVLRCGDLRRPMLCTKCAYDHVFGT	5032
10	Query: 2882 DHKFILAITPYVCNASCGGVSDVKLYLGGLNYYCTNHKPQLSFPLCSAGNI FGLYKNSA DHKFILAITPYVCN SGC V+DV KLYLGGLNYYC +HKP LSFPLCSAGN+FGLYK+SA	3061
Sbjct: 5033	DHKFILAITPYVCNTSGCNVNDVTKLLYLGGLNYYCVDHKPHLSFPLCSAGNVFGLYKSSA	5092
15	Query: 3062 TGSLDVEVFNRILATS DWDVRDYKLANDVKDTLRLFAAETI KAKEESVKSSYAFATLKEV GS+D++VFN+L+TSDW+D+RDYKLAND K++LRLFAAET+KAKEESVKSSY+A+T+LKE+ Sbjct: 5093 LGSMIDDFVNKLSTSDWSIDRDYKLANDAKESLRLFAAETVKAKEESVKSSYAYATLKEI	3241
20	Query: 3242 VGPKELLLSWESGKVKPPLNRNSVFTCFQISKDSKFQI GEF FIEKVEYGSDTVTKSTVT VGPKELLLL WESGK KPPLNRNSVFTCFQI+KDSKFQ+GEF+FEKV+YGSDTVTKST T Sbjct: 5153 VGPKELLLLWESGKAKPPLNRNSVFTCFQITKDSKFQVGEFVFEKVDYGSDTVTKSTAT	3421
25	Query: 3422 TKLVPGMIFVLTSHNVQPLRAPTIANQEKYSSIIYKLHPAFNVSDAYANLVPYYQLIGKQK TKLVPGM+F+LTSHNV PLRAPT+ANQEKYS+IYKLHP+FNVSDAYANLVPYYQLIGKQ+ Sbjct: 5213 TKLVPGMLFILTSHNVAPLRAPTMANQEKYSTIYKLHPSPNVSDAYANLVPYYQLIGKQR	3601
Sbjct: 5213	ITTIQGPPGSGKSHCSIGLGLYYPGARIVFVACAAVDSLCAKAMTVYSIDKCTRIIPA	5272
30	Query: 3602 ITTIQGPPGSGKSHCSIGLGLYYPGARIVFVACAAVDSLCAKAMTVYSIDKCTRIIPA ITTIQGPPGSGKSHCSIG+G+YYPGARIVF AC+AAAVDSLCAKA+T YS+DKCTRIIPA	3781
Sbjct: 5273	ITTIQGPPGSGKSHCSIGIGVYYPGARIVFTACSHAAVDSLCAKAVTAYSVDKCTRIIPA	5332
35	Query: 3782 RARVECYSGFPKNNTSAQYIFSTVNALPECNADIVVVDEVS MCTNYDL SVINQR LSYKHI RARVECYSGFPKNN SAQY+FSTVNALPE NADIVVVDEVS MCTNYDL SVINQR+SYKHI	3961
Sbjct: 5333	RARVECYSGFPKNNNSAQYVFSTVNALPEVNADIVVVDEVS MCTNYDL SVINQR ISYKHI	5392
40	Query: 3962 VYVGDPQQLPAPRVMITKGMEPV DYNVVTQRMCAIGPDVFLHKCYRCPAEIVNTVSELV VYVGDPQQLPAPRVI+I+KGMEP+DYNVVTQRMCAIGPDVFLHKCYRCPAEIVNTVSELV	4141
Sbjct: 5393	VYVGDPQQLPAPRVLISKGVMEPIDYNVVTQRMCAIGPDVFLHKCYRCPAEIVNTVSELV	5452
45	Query: 4142 YENKFVPVKPASKQCFKIFFKGNVQVDNGSSINRKQLEIVKLFLVKNPSWSKAVFISPYN YENKFVPVK ASKQCFKIF +G+VQVDNGSSINR+QL++VK F+ KN +WSKAVFISPYN	4321
Sbjct: 5453	YENKFVPVK EASKQCFKIFERGSVQVDNGSSINRRQLDVVKRPIHKNSTWSKAVFISPYN	5512
50	Query: 4322 SQNYVASRFLGLQIQTVDSSQGSEYDVVIYAQTSDTAHACNVRFNVAITRAKKGIFCVM SQNYVA+R LGLQ QTVDSS+QGSEYDVVI+AQTSDTAHACN NRFNVAITRAKKGIFC+M	4501
Sbjct: 5513	SQNYVAARLLGLQTQTVDSAQGSEYDVVIYAQTSDTAHACNANRFNVAITRAKKGIFCIM	5572
55	Query: 4502 CDKTLFDSLKF FEIKA DHLHSQVCGLFKNC TRTPNLPPTHAHTFLSLSQDFKTTGDLA D+TLEDF+LKF FEI DL S CGLFK+C R P++LPP+HA T+LSLSD+FKT+GDLA	4681
Sbjct: 5573	SDRTLFDALKF FEI TMTDLQSESSCGLFKDCARNP IDLPPSHATTYLSLSDRFKTSGDLA	5632
60	Query: 4682 VQIGSNNVCTYEHVISFMGFRFDISI PGSHSLFCTRDFAIRNVRGWLGM DVESA HVGCDN VQIG+NNVCTYEHVIS+MGFRFD+S+PGSHSLFCTRDFA+R+VRGWLGM DV AHV GDN	4861
Sbjct: 5633	VQIGNNNVCTYEHVISYMGFRFDVSMPGSHSLFCTRDFAIRNVRGWLGM DVEGA HVTGDN	5692
65	Query: 4862 IGTNVPLQVGF SNGVN FVVQTEGV STNF GVDV1KPVCAKSPPGEQFRHLPFLRKQGPWL +GTNVPLQVGF SNGV+FV Q EGV TN G V+KPV A++PPGEQF H+VP LRKGQGPW	5041
Sbjct: 5693	VGTNVPLQVGF SNGVDFVAQPEGCVL TNGSVV KPV RARAPP GQF THIVPLLRKGQGPWS	5752
70	Query: 5042 IVRRRIVQMI S DYL S NL SDIL FV L WAG S LE LTTM RYF V KIG PI K YC CG NSAT C YNS VS ++R+RIVQMI+D+L+ SD+L FV L WAG LE LTTM RYF V KIG +K+C CG AT C YNS VS	5221
Sbjct: 5753	VLRK RIVQMI ADFLAGSSDVL FV L WAG G LE LTTM RYF V KIG AVKHCQCGTVAT C YNS VS	5812
75	Query: 5222 NEYCCFKHALGCDYVNPYAFDIQOWGYVGSLS 5320	
Sbjct: 5813	N+YCCFKHALGCDYVNPYDIOQOWGYVGSLS 5845	

## 65 5. Sequence E

6143 nucleotides; 3' end of Replicase and 5' end of Spike

TCTGGAATTGTAATGTTgATATGTATCCAGAATTTCATAATTGTGTGTCGCTTGACACACGTACTCGTTCTGTTT  
TTAATTAGAAGGTGTTAATGGGGTTCTCTTATGTTAACAAACATGCGTTCATACACCAGCATATGATAAAC  
GTGCTTTGTTAAATTAAACCTATGCCCTTTTTACTTTGATGACAGTGATTGTGATGTTGCAAGAACAAAG  
TTAATTATGTACCCCTTCGCGCTAGTAGTTGTGTTACCGTTGTAATATAGGTGGTGCTGTTGTTCAAAACATG  
CAAATTGTTATCAAAAATAGTTGAGGCATATAACATTACACAGGCTGGTTTAACATTGGGTACCAACATA  
GTTTGATGTTATAATTGTTGGCAAAATTTTATTGAAACTAATTACAAAGTCTTGAAAATATAGCATTTAATG  
TTGTAaaaaaaGGGTGTTTACTGGTGTGATGGTAGTTACCTGTTGAGTTACGTTAACGACAAAGTTTGTTG  
GCTATGGCGATGTTGACAATTGGTTTACAATAAACAAACATTGCCACTAATGTTGTTGTTGAATTGTTTG

CAAAACGAAAAATGGGTTAACACCACATTGCTATTCTAAAAATCTGGTGTGCTACATATAAAATTG  
 TTTTATGGATTATGAAGCTGAAAGACCTTTACCTCATATACTAAGAGTGTATGTAATACACTGATTAAATG  
 AGGATGTTGTTGTTGACAATAGTATTCAAGGGTTCGTATGAGCGTTTACGTTACTACGAACGCTGTT  
 TATTTCTACTGTTGTCATTAACACCTATAAAGTGAATTGGTATGTTGAATGGTATGCCAGTT  
 5 CTTCTATTAAGAGTGATAAAGGTGTTGAAAAATTAGTTAATTGGTACAYATATGTTGCTAAAATGGTCAATTTC  
 AAGATCATTATGATGGTTTACACTCAAGGTAGGAATTATCAGACTTACACCAAGAAGTGTATGGAGTATG  
 ATTTCTTAACATGGATATGGTGTATTAAATAAAATGGTCTGAGGATTAAATTGGAACATGTTGTT  
 ATGGTGTATGTTCAAAAACATCAATTAGGCTTCTCATTGTTGATATCACAGGTTAGGTTAGTAAATGGGTG  
 10 TTTGAAAGCTGATGATTGGTCACTGCTCTGACACAAACTTGGAGGTGCTGACTGTTACTTATCTTAATGAAC  
 GTGTTAAGCTGTTAAATACACTCAGCTTGTCAATACCTAAATAGCACTACATGTGCGTACCTCATAATG  
 15 GTGTTTGCACATGGTGTGGTTCTGACAAAGGTGTCACACTGTTAAAAGCTGGTAAAGATA  
 CTGATGCAATAATCATTGATAATGATATCAATGATTATGTTAGGCTGAGATTGATACAGGTGATTG  
 CTACTGTTACCTGAAAGATAAGGTTGACTTACTTATCTGATATGTTAGGCTGTTAAGGTCCTAGGGTATAA  
 GTGAAAAGCTCTAAAGATGGTTTTACTATCTTAATGGTGTATTAGAGAAAAATTAGCTATTGGTGT  
 GTGTTGCCATTAAAGTACAGAATATAGTGGATAAGTATCTTATGAATTAAACAAAGATTGCTTTGG  
 20 CTTTGTCTGACGTCTGTTAATACATCCTCTCAGAACGCTTTCTATTGGTATTAAATTATTAGGT  
 TTCAAGGTCTTTATAGCTGGTAACACTGTTCATGCTAATTATATATTGGCTAATTCTACTATTGCTT  
 TGTCTACAAATTCAAGTTAGATTAAAGTGAATGTAACATAAGGCACACTGTTGTTACACTAAAG  
 ATAGTGATGTAATGATATGGTTTGAGTTAGGTTGAGGTTGTTACGTTAAGTGGCCGTTTG  
 25 GTGGTTTAGTAACTTCTGATGTTCAACTAAATGAAACATTCTTCTGATTGGCTTATTTGCTT  
 CTTTCTACATGTAACAGTAATGCTAGTATTCTATGTTACAATTAGGTGTTCTGATAACTCTCAACTATTG  
 CACAGGTTGTCAGTCATTGGATTGCTAATCAGAGTACATCTAGTTACCCAGCCAACGGCTTTCTA  
 TATTGATGTTGTAACACCCAGTAGTCCTGACTCCATAGGGTTATTATGATGCTAACCAAGTATTATTTA  
 TCTCTAAATAAAACATTTAAATGCTCTGACTCTGAAAGATTGTAAGTTGGAAACACTCTTTGATT  
 TTTAAGTAATGTTCTACTCTCATGATTGATAGTTAATTGTCATTACAGAACAGTTAGGTGTCCTTTGG  
 30 CATAACTATATCGGGTGAACACTGTACGTTGCATTATATAATGCAACTCGTACTTTTATGTCGGGGCGCTTA  
 TAAACTTAACTAAACTTAGTGTAAATGTTACTTATGTAATCCTGTTTTAGTGTGTTCAATGCCACATTAC  
 TGTTAATGTCACCACACTTAATGGCGTATAAGTTAACTACACTGTTGATGATTGTAATGGTTACCTGATAA  
 CATATTCTGTTCAACAGGATGGCGCATTCTTAATGGTTTCTTAAATAATTGGTTTGTAACTAATGG  
 TTCCACATTAGGGACGGGCTCTAGACTTATCAACACACTCCGTTAACTTGTATGGCCTGTACCTGGTCT  
 35 TAAATCTCACTGGTTTGTGTTATTAAATGCCACTGGTTCTGATGTTAATTGTAACGGCTATCAACATAATT  
 TGTTGCTGATGTTATGCTTACAATCTTAACTCAGTGTAACTTGTGCTAATTCTGTGACAACTTAAAGAGTGGTT  
 TTTAAAACCTTACAGTACGATGTTGTTATTGTTAGTAATTCTTCTCAGGTGTTCTGACACCACAAATTAC  
 TTTGGCCCTCCTCTCAACCTTAACTGTTATAAACAGTACTATCAACACTACTCATGTTAGCACTTTGT  
 GGGTATTACCAACCACTGTGCGTGAATTGTTGCTAGAACACTGGTAGTTTATTAATTGTTAAAGTA  
 40 TTTCGATTGGGTTCTAGAAGCTGTCATTAAATGTCACGACTGGTAGTGCACAGGTTTATGCACTTCAGTGT  
 ATTGCTACTTTGGTATGTTGGTTAATGTTAGTGTGCAACTAACATTCAAAACTTACTTTATGCGATTCTCC  
 ATTGAAAAGCTGAGTGTGAGGACTTGCAGTTGGATTGCAAGATGGTTTTATTCTGCAAATTTCCTGATGA  
 TAAATGTTGCTGAGACTTATGTTGACTCCCAATTATTCAACATACGGACATAATTAACTGCAACTGC  
 ATCTTTGGGGTTCTGTTATGTTAAACCACGCCAGGTTAATATATCTTAATGGTAACACACTTCAGTGT  
 45 TGTAGAACATCTCATTTCAATTAGGTATATTATAACCGCTTAAGAGTGGTTACCCAGGTGACTCTCATG  
 GCATATTATTAAGAGTGGCACTTGTCCATTCTTCTAAGTTAAATAATTCTCAAAGGTTAAAGACTAT  
 TTGTTCTCAACCGTCAAGTGGCTGGTAGTTGTAATTCTCAGGCAACCTGGCATTACACTCTTATAC  
 TATTGTTGGGCTTTGTTATGTTACTTGTGCTGAAGGTAATTCCATTACTGGTTGACCTTATCCTGCTGGTAT  
 TCGTAGTTAGTAATTGTTAAATGTTACCAAAATATAATTATGTTATGTTGTTACTGGAAATTAT  
 50 ACGTTCTCAACACAGTCACTTGTGGGGTATTACATATGTTCTAACTCTGGTAATTACTGGTTTAAAAA  
 TGTTTCCACTGGTAACATTATTGTCACGGCTGTTAATGAGCTTACCTGCAACTAACATTCCATTCT  
 TATTGGGCCATTGACCGCTGTTAATGAGTCTAGATATGGCTGCAAAACTTACTACAGTTACCTAATT  
 TGTAGTAATGGGGTAACAATTGCAACTACGGCTGTTATGATTATTCTAATTGGTATTGTCATGGTIC  
 TTTAATTCTGTGCTGGCTGTAATTCTAGTGTAAATTGTTACTTGTGCTGTTATTCTAATTGTT  
 55 CTCTAACTGGACTACTTCAGTCAAGTGTGAGTACCTCCTAAATTACTAGTACTCCAAATGTTGTTGCTAC  
 TTATGTTGTTAATGGTAACCCCTCGTTGTAAGAATCTACTAAGCAGTAACTTGTGTTGCTGTT  
 TGCCCTACGACTTGTGCTCATTTGAAACATTGATGTTAGTAGTACTGCTAACTTCTGCTGTTAAGAAGA  
 TTTGGCTAATGTTACTAGTTGGAGATTATAACCTTCTAGTGTGTTACCTCAGAGAAACATTCAAGCCG  
 TATAGCAGGAGCTAGTGTCTTGGAGATTGTTGTTAGCAAAAGTGTGTTACATCTGGTTGGGACTGTTGATGT  
 60 TGACTATAAGCTTGTACTAAAGGTCTTCTATTGTCACGGCTGCTGCTGCTCAGTACTACAAATGGCATAATGGT  
 TTTGCCAGGTGTTGCTGATGCTGAACGTATGCCATTGTCACACAGGTTCTTATAGGTGGCATTGGCTCGGAGG  
 TCTTACATCAGCAGCCCATACCTTTCTGGCACTGCAAGCAGACTAACTATGTTGCTTACAAACTG  
 TGTGCTCAAGAAAATCAGAAAATTGGCTGATCTTAATAAGCTATTAAATAATTGTTGCTTCTTTAG  
 TAGCGTTAATGATGCTTACACACTGCAAGAGGCTATACTACATGTTACTATTGCACTAATAAGGTCAGGA  
 65 TGTGTTAATCAACAGGGTAGTGTCTTAACCATCTCAGTCACTTCAACATTGAGACATAATTTCAGGCCATTCTAA  
 TTCAATTCTGCTATTGACGGCTTGTGATTCAATTCAAGCCGATCAACAAAGTGTGACAGATTAAATTACTGGACG  
 GCTGCAAGCTTGTGATGCAATTGCTTCCAAAGTTGTTGAAATAAAATACTGAAAGTTCGTTGGCTCAGCGCTAGC  
 ACAGCAGAAGATTAAATGAATGTCAGTCACAACTTAATAGATATGGTTTGTGGCAATGGCAACTCACATCTT

5 TTCAATCGTCAACTCAGCTCCAGATGGTTGCTTTCTCATACTGTTTGCTGCCAAGTGAATTACAAGAATG  
 AAAGGCGTGGTCTGGTATCTGTGTTGATGGCATTTATGGCTATGTTCTGCGTCAACCTAACCTGGTTCTTATT  
 TGATAATGGGTCTTCGTGAACTTCAGGGCATGTTCAACCTCGTTACCTGTTTGCTGATTGTGC  
 AATATATAATTGTAATGTTACTTGTAAACATATCTCGTGTGAGTTACATACGTACACTGACTACGTTG  
 5 TGTTAATAAAACATTACAAGAGTTGCACAAACTACCAAGTATGTTAACGCTAATTGACTTGACTCCTT  
 TAATTAAACATATCTAATTGAGTTCTGAGTTGAAGCAACTCGAAGCTAAACTGCTACGAATCAGC

Hypothesised ORFs

10 >-out: 3 to 2357: Frame 3 785 aa  
 WNCNVDMYPEFSIVCRFDTRRSVFNLEGVNGGSLYVNKHAFHTPAYDKRAFVKLKPMFFYFDDSDCDVVQEQ  
 NYVPLRASSCVTRCNIGGAVCSKHANLYQKYVEAYNTFTQAGFNIWVPHSFVDVNLWQIFIETNLQSLNIAFN  
 VKKGCFGTGVDGEPEVAVVNDKVFVRYGVDVNLVFTNKTTLPNVAFELFAKRKGMLTPPLSILKNLGVVATYKF  
 LWDYEAEERPFTSYTKSVCKYTFNEDCVCFDNIQGSYERFTLTNAVLFSTVVVIKNLTPIKLNFGMNGMPV  
 SIKSDKGVEKLVNVNXYVRKNGQFQDHYDFYQTQGRNLSDFTPRSDMEYDFLNMDMGVFINKYGLDFNFEHVV  
 15 GDVSKTTLGGHLIISQFRLSKMGVLKADDFTASDTTLRRCVTYLVNLSSKVCTYMDLLLDFVTILKSLD  
 GVISKVHEVIIDNPKYRWMLWCKDNHLSTFYPQLQSAEWKCGYAMPQIYKLQXMCLEPCNLYNYGAGIKLPSGI  
 LNVVKYTQLCQYLNSTTMCPVPHNMRLVHYGAGSDKGVAPGTTVLKRWLPPDAIIIDNDINDYVSDADFSITGDC  
 TVYLEDKF DLLISDMYDGRIKFCDEGENSKDGFETYLNGBIREKLAIGGSVAIKITEYSWNKLYELIQRFAFW  
 20 LFCTSVNTSSSEAFLIGINYLGFIQGPFIAVNTVHANYIFWRNSTIMSLSYNSVLDLSKFECKHKATVVVTLK  
 SDVNDMVLSLIKSGRLLLRLNSGRFGFSNHLVSTK  
 >-out: 277 to 438: Frame 1 54 aa  
 VVLFVQNMQICIKNMLRHIIHLHRLVLTFGYHIVLMIICGKFLKLKLYKVLKI  
 >-out: 457 to 618: Frame 1 54 aa  
 KKGVLLVLMVSYLLQLLTTKFLFAMAMLTTWFLQIKQHCLIMLLLNLCLQNEKWV  
 25 >-out: 622 to 852: Frame 1 77 aa  
 HHHCLFSKILVLLHINLFLYQGIMKLKDLLPHILRVYVNTLILMRFVVFVLTIVFRVRMSVLRLRTLFYFLLLS  
 KI  
 >-out: 937 to 1149: Frame 1 71 aa  
 LIGTXMFVKMVNFKIIIMMVFTLKVGIYQTLHQEVISMIIFTWIWVFLINMVLRILILNMLYVMVFQKLH  
 30 >-out: 1387 to 1572: Frame 1 62 aa  
 IINLIGGCCGVKITTCPLFIHSCSLLNGSVVMLCHKFISFXNCVWNLVIIYIIMVVLVLSCLVV  
 >-out: 1738 to 1935: Frame 1 66 aa  
 SLIMISMIMLVMQILALQVIVLFLKISLTYLFLICMVMVELNFMVKTSLSKMVFLLILMVILLEKN  
 >-out: 2357 to 6142: Frame 2 1262 aa  
 35 MKLFLILLILPLVSCFSTCNSNASISMLQLGVPDNSSTIVTGLLPVHWICANQSTSSYPANGFFYIDVGKHRSAJ  
 ALHSGYYDANQYYIYLTKIHLNAPVTLKICKFGNTSFDFLSNVSTSHDCIVVNLSTFEQLGVPLGITISGETVRI  
 HLYNATRTFYVPAAYKLTKLSVKCYFSESCVFVSVNATITVNVITLNGRIVNYTCDDCNGYTDNIFSVQDGR  
 PNGFPFNWFLLTNGSTLVDGVSRLYQPLRLTCLWPVPLKSSSTGFVYFNATGSDVNCNGYQHNSVADVMRYNL  
 LSANSVDMNLKSGVIVFKTLQYDVLFYCSNSSSGVLDTTIPFGPSSQPYCFCINSTINTTHVSTFVGILPPTVRE  
 40 VVARTGQFYINGFKYFDLGFTEAVNFNVTTASATDFTWVAFATFVDFVLVNSATNIQNLLYCDSPFEKLQCEHL  
 FGLQDGFFYANFLDDNVLPETYVALPIYYQHTDINFTATASFGGSCYVCKPRQVNISLNGNTSVCRTSHFSIR  
 IYNRVKSGSPGDSSWHIYLKSGTCPFSKLNQFKFTICFSTVEPGSCNFPLEATWHYTSYTVI GALYTVW  
 EGNSITGVYPVSGIREFSNLVNNCTKNIYDVGVTGIIRSSNQSLAGGITYVSNSGNLLGFKNVSTGNIFIV  
 45 PCNQPDQAVYQQSIIGAMTAVNESRYGLQNLQLPNFYVSNNGNNCTAVMIYSNFGICADGSLIPVRPRNS  
 DNGISAIITANLISIPSNWTTSVQVEYLQITSTPIVVDCATYVCNGNPRCKNLLKQYTSACKTIEDALRLSAHLET  
 NDVSSMLTFDSNAFSLANVTSVQVEYLQITSTPIVVDCATYVCNGNPRCKNLLKQYTSACKTIEDALRLSAHLET  
 IADLACAQYNGIMVLPGVADAERMAMYTGSLIGGMVLGGLTSAAAIPFSLALQARLNQVALQTDVLQENQKIL  
 ASFNKAIIINNIVASFSSVNDAITHAEEAHTVTTALNKIQDVNVQQGSALNHLTSQRLHNFOAISNSIHA  
 50 SIQADQQVDRLLITGRLLAALNAFVSQVLNKYTEVRGSRLLAQKQINECVKSQSNSRYGFCGNGTHIFSI  
 LFLHTVLLPTDVKNVKAWSGICVCDGIYGYVLRQPNLVLYSDNGVFRVTSRVMFQPRLPVLSDFVQIYNC  
 ISRVELHTVIPDYDVNKTLQEFQANLPKVKPNFDLTPFNLTYLNLSSELKQLEAKTATNQ  
 >-out: 2448 to 2645: Frame 3 66 aa  
 VFLITLQLLSQVCCQSIGFVLIRVHLVTQPTAFSISILMLVNTVPLHSIVVIMMLTSIIFISLIKYI  
 >-out: 2781 to 2954: Frame 3 58 aa  
 55 LYRVKLYVCIYIMQLVLFCRPLINLLNLVNVTLVNPVFLVLSMPPLLMSPHLMAV  
 >-out: 3126 to 3296: Frame 3 57 aa  
 LVYGLYLVNLQLVLFILMPLVLMILVTAINTIILLMLCVTILTSVLLWTLRVVL  
 >-out: 3546 to 3806: Frame 3 87 aa  
 KLSILMSRLLVQPIFGRLHLLLLMFWLMLVQLTFKTYFIAILHLKSCSVSTCSLDCKMVFILQIFLMIMFCLRL  
 60 MLHSPFTIINIRT  
 >-out: 3810 to 3986: Frame 3 59 aa  
 TLLQLHLLVVLVMFVNHARLIYLLMVTIQCVLLEHLIFQLGIFITALRVVHQVTLHGIFI  
 >-out: 4026 to 4217: Frame 3 64 aa  
 IIIFKSLRLFVSPSKCLVVVIFHLKPPGIFTLLLVLCLMGLKVIPLLVYLILSLVFVSLVI  
 65 >-out: 4227 to 4376: Frame 3 50 aa  
 IIIVPNIIIFMIMLVLELYVLQTSVHLLVVIHMFLTLVVIYLVLKMFPLVTFLL  
 >-out: 5157 to 5447: Frame 3 97 aa

VAWCSEVLHQOPPYLFLWHCKHDLTMLLYKLMCFKKIRKFWLHHLIRLLIILLLLLVALMMLLHILQRLYILLLL  
 HLIRFRMILLINRVVLLTISLHN  
 >-out: 5625 to 5774: Frame 3 50 aa  
 HSRRIMNVSSHNLIDMVFVAMALTSFQSSTQLQMVCFFFILFCCQLITRM  
 5 >-out: 5874 to 6065: Frame 3 64 aa  
 LPGSCFNLVYLFCLILCKYIIVMLLLTLYVSSYILSYLTLMLIKHYKSLHKTYQSMLSLILT

### Alignment

10 >gi|12175747|ref|NP\_073549.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 >gi|30179827|sp|Q05002|R1AB CVH22 Replicase polyprotein 1ab (pp1ab) (ORF1ab polyprotein) [Includes:  
 Replicase polyprotein 1a (pp1a) (ORF1a) [Contains: p9;  
 p87; p195 (Papain-like proteinases 1/2)  
 15 (PL1-PRO/PL2-PRO); Peptide HD2; 3C-like proteinase  
 (3CL-PRO) (3CLp) (M-PRO) (p34); Unknown protein 1; p5;  
 p23; p12; Growth factor-like peptide (GFL) (p16);  
 RNA-directed RNA polymerase (RdRp) (Pol) (p100); Helicase  
 (Hel) (p66) (p66-HEL); Unknown protein 2; p41; Unknown  
 20 protein 3]  
 >gi|12082740|gb|AAG48591.1| replicase polyprotein 1ab [Human coronavirus 229E]  
 Length = 6758

Score = 1332 bits (3448), Expect = 0.0  
 25 Identities = 630/789 (79%), Positives = 695/789 (88%), Gaps = 4/789 (0%)  
 Frame = +8

Query: 3 WNCNVDMPPEFSIVCRFDTRTRSVNLEGVNGSLYVNKHAFHTPAYDKRAFKVLKPMPF 182  
 WNCNVDMPPEFSIVCRFDTRTRSVNLEGVNGSLYVNKHAFHTPAYDKRAFKVLKPMPF 182  
 30 Sbjct: 5970 WNCNVDMPPEFSIVCRFDTRTRSVNLEGVNGSLYVNKHAFHTPAYDKRAFKVLKPMPF 6029

Query: 183 FYFDDSDCDVVQEQQVNYVPLRASSCVTRCNIGGAVCSKHANLYQKQYVEAYNTFTQAGFNI 362  
 FY+DD C+VV +QVNYVPLRA++C+T+CNIGGAVCSKHANLY+ YVE+YN FTQAGFNI  
 35 Sbjct: 6030 FYYDDGSCEVVHDQVNYVPLRATNCITKCNIGGAVCSKHANLYRAYVESYNIFTQAGFNI 6089

Query: 363 WVPHSFDVYNLWQIFIETNLQSLENIAFNVVKKGCFGTGVDGELPVAVVNDKVFVRYGDVD 542  
 WVP +FD YNLWQ F E NLQ LENIAFNVV KG F G DGEELPVA+ DKVEVR G+ D  
 Sbjct: 6090 WVPPTTFDCYNLWQTFTEVNLQGLENIAFNVVNKGSFVGADGELPVAISGDKVEVRDGNTD 6149

40 Query: 543 NLVFTNKTTLPTNVAFELFAKRKMGLTPPLSILKNLGVVATYKFVLWDYEAERPFTSYTK 722  
 NLVFT NKT+LPTN+AFELFAKRK+GLTPPLSILKNLGVVATYKFVLWDYEAERP TS+TK  
 Sbjct: 6150 NLVFTNKTTLPTNIAFELFAKRKVGLTPPLSILKNLGVVATYKFVLWDYEAERPFTSYTK 6209

45 Query: 723 SVCKYTDNFEDVCVCFDNSIQGGSYERFTLTTNAVLFSTVVIK---NLTPIKLNFGMLNG 890  
 SVC YTDFAEDVCVCFDNSIQGGSYERFTLTTNAVLFSTVVIK---NLTPIKLNFGMLNG  
 Sbjct: 6210 SVCGYTDFAEDVCVCFDNSIQGGSYERFTLTTNAVLFSTVVIK---NLTPIKLNFGMLNG 6269

50 Query: 891 MPVSSIKSDKGVEKLVNWYXYVORKNGQFQDHYDFYQGRNLSDFTPRSDMNEYDFLNMDM 1070  
 +++++KS+ G K +NW+ YVRK+G+ DHYDFYQGRNL DF PRS ME DFLNMD+  
 Sbjct: 6270 NAIATVKSEDGNITKNNINWVFVYVORKDGKPVDHYDFYQGRNLQDFLPRSTMEEDFLNMDI 6329

55 Query: 1071 GVFINKYGLEDFNFEHVVYGDVSKTTLGGHLHLLISQFRLSKMGVLKADDFTVASDTTLRC 1250  
 GVFINKYGLEDFNFEHVVYGDVSKTTLGGHLHLLISQFRLSKMGVLKADDFTVASDTTLRC  
 Sbjct: 6330 GVFINKYGLEDFNFEHVVYGDVSKTTLGGHLHLLISQFRLSKMGVLKADDFTVASDTTLRC 6389

60 Query: 1251 CTVTYLNESSKVVCTYMDLDDDFVTLKSLDLGVISKVKHEVIIIDNKPYRWMLWCKDNH 1430  
 CTVTYLNESSKVVCTYMDLDDDFVTLKSLDLGVISKVKHEVIIIDNKPYRWMLWCKDNH  
 Sbjct: 6390 CTVTYLNESSKVVCTYMDLDDDFVTLKSLDLGVISKVKHEVIIIDNKPYRWMLWCKDNH 6449

65 Query: 1431 LSTFYPQLQSAEWKCGYAMPQIYKLQRMCLEPCNLNYGAGIQLPSGIMLNVVKYTQLCQ 1610  
 ++TYPQLQSAEWKCGYAMPQIYKLQRMCLEPCNLNYGAGIQLPSGIMLNVVKYTQLCQ  
 Sbjct: 6450 VATFYPQLQSAEWKCGYSMPGIYKLQRMCLEPCNLNYGAGIQLPSGIMFNVVKYTQLCQ 6509

70 Query: 1611 YLNSTTMCVPHNMRLHYGAGSDKGVAPGTTVLKRWLPPXXXXXXXXXXVSDADFSIT 1790  
 Y NSTT+CVPHNMRLHYGAGSDKGVAPGTTVLKRWLPPXXXXXXXXXXVSDADFSIT  
 Sbjct: 6510 YFNSTTLCVPHNMRLHYLHGAGSDYGVAPGTTVLKRWLPHDAIVVNDVVDYVSDADFSIT 6569

Query: 1791 GDCATVYLEDKFDLISDMYDGRIKFCDFGENVSKDGFFTYLNGVIREKLAIGGSVAIKIT 1970  
 GDCATVYLEDKFDLISDMYDGRIKFCDFGENVSKDGFFTYLNGVIREKLAIGGSVAIKIT  
 70 Sbjct: 6570 GDCATVYLEDKFDLISDMYDGRTKAIDGENVSKEGFFTYINGFICEKLAIGGSIAIKVT 6629

Query: 1971 EYSWNKYLVELIQRFAFWTLFCTSNTSSSEAFILGINYLGDFIGQPFIAGNVTWANYIF 2150  
 EYSWNKYLVELIQRFAFWTLFCTSNTSSSEAFILGINYLGDFIGQPFIAGNVTWANYIF

Sbjct: 6630 EYSWNKKLYELVQRFSFWTMFCTSVNTSSSEAFVVGINYLGDFAQGPFIDGNIIHANYVF 6689  
 Query: 2151 WRNSTIMSLSYNSVLDLSKFECKHKATVVVTLKDSDVNDMVLSLIKSGRLLLNSGRFGG 2330  
 5 Sbjct: 6690 WRNSTVMSLSYNSVLDLSKPNCKHKATVVVQLKDSDINEMVLSLVRSGKLLVRGNGKCLS 6749  
 Query: 2331 FSNHLVSTK 2357  
 FSNHLVSTK  
 Sbjct: 6750 FSNHLVSTK 6758  
 10  
 >gi|13604832|gb|AAK32188.1| spike glycoprotein [Human coronavirus 229E]  
 Length = 1173  
 15 Score = 1891 bits (3600), Expect = 0.0  
 Identities = 682/1069 (63%), Positives = 833/1069 (77%), Gaps = 7/1069 (0%)  
 Frame = +2  
 20 Query: 2948 GRIVNYTCDDCNGYTDNIFSVQQDGRIPNGFPFNNWFLLTNGSTLVDGVSRVLYQPLRLT 3127  
 G +Y+VC+ C GY++N+F+V+ G IP+ F FNNWFLLTN S++VDGV R +QPL L  
 Sbjct: 21 GLNTSYSVCNGCVGYSENVFAVESGGYIPSDFAFNNWFLLTNTSSVVDGVVRSFQPLLLN 80  
 25 Query: 3128 CLWPVPGLKSSSTGFVYFNATGSDVNCGYQHNSVADVMRYNLLNLSANSVDNLKSGVIVFK 3307  
 CLW V GL+ +TGFVYFN TG +C G+ + +DV+RYNLN +NL+ G I+FK  
 Sbjct: 81 CLWSVSGLRTTGFVYFNGTGRG-DCKGFSSDVLSDVIRYNLNFE---ENLRRGTILFK 135  
 30 Query: 3308 TLQYDVLFYCSNSSGVLDTTIPFGPSSQPYCFINSTINTTHVSTFVGILPPTVREIVV 3487  
 T V+FYC+N++ D IPFG +YCF+N+TI S FVG LP TVRE V+  
 Sbjct: 136 TSYGVVVFYCTNNILVSGDAHIFPFGTVLGNFYCFVNTTIGNETTSAFVGALPKTVREFVI 195  
 Query: 3488 ARTGQFYINGFKYFDLGFIEAVNFNVTASATDFWTVAFAFFVVDVLVNVSATNIQNLLYC 3667  
 +RTG FYING++YF LG +EAVNFNVTAA TDF+TVA A++ DVLVNV S T+I N++YC  
 Sbjct: 196 SRTGHFYINGYRYFTLGNVEAVNFNVTAAETTDFFTVALASYADVLVNVSQTSIANIYC 255  
 35 Query: 3668 DSPFEKLQCIEHLQFGLQDGFSANFLDDNVLPETYVALPIYYQHTDINFAT---ASF GG 3838  
 +S +L+C+ L F + DGFYS .+ + LP + V+LP+Y++HT I S GG  
 Sbjct: 256 NSVINRLRCQDQLSFVDPDGYSTSPIQSVLPVSI VSLPVYHKTIFIVLYVDFKPQSGGG 315  
 40 Query: 3839 SCYVCKPRQVNISL-NGNTS---VCVRTSHFSIRYIYNRVKSGSPGDSSWHIYLKSGTCP 4006  
 C+ C P VNI+L N N + +CV TSHF+ +Y+ G W + +G CP  
 Sbjct: 316 KCFNCYPAGVNITLANFNETKGPLCVDTSHFTKYVAVYANVGR----WSASINTGNCP 370  
 45 Query: 4007 FSFSKLNQFQFKTICFSTVEVPGSCNFPLEATWHYTSYTIIVGALYVTWSEGNSTITGPY 4186  
 FSF K+NNF KF ++CFS ++PG C P+ A W Y+ Y +G+LYV+WS+G+ ITGVP  
 Sbjct: 371 FSFGKVNNFVKFGSVCFSLKDIPGGCAMPIVANWAYSKYTTIGSLYVWSWSDGDGITGPQ 430  
 50 Query: 4187 PVSGIREFSNLVNNCTKYNIYDVGTCIIRSSNQSLAGGITVVSNSGNLLGPKNVSTGN 4366  
 PV G+ F N+ L+ CTKYNIYD G G+IR SN + CITY S SGNLGFK+V+ G  
 Sbjct: 431 PVEGVSSFMNVLDKCTKYNIYDVGVGIRVSNDTFLNGITYTSTSGNLLGFKDVTKGT 490  
 Query: 4367 IFIVTPCNQPDQVAVYQQSIIGAMTAVNESRYGLQNLLQLPNFYYVSNGGNCTTAVMIY 4546  
 I+ +TPCN PDO+ VYQQ+++CAM + N + YG N+++LP F+Y SNG NCT AV+ Y  
 Sbjct: 491 IYSITPCNPPDQLVYVYQQAVVGAMLSENFTSYGSNVVLPKFFYASNGTYNCTDAVLTY 550  
 55 Query: 4547 SNFGICADGSLIPVPRPRNSSDNGISAIITANLSIPSNWTSVQVEYLQITSTPIVVDCAT 4726  
 S+FG+CADGS+I V+PRN S + +SAI+TANLSIPSNWTSVQVEYLQITSTPIVVDCT+S  
 Sbjct: 551 SSFGVCADGSIIAVQPRNVSYDSVSAITANLSIPSNWTSVQVEYLQITSTPIVVDCT 610  
 60 Query: 4727 YVCNGNPRCKNLLKQYTSACKTIEDALRLSAHLENDVSSMLTFDSNAFLANVTSGDY 4906  
 YVCNGN RC LLKQYTSACKTIEDALR SA LE+ DVS MLTFD AF+LANV+SGDY  
 Sbjct: 611 YVCNGNVRCVELLKQYTSACKTIEDALRNSARLESADVSEMLTFDKKAFTLANVSSFGDY 670  
 65 Query: 4907 NLSSVLPQRNIHSSRIAGRSALEDDLLFSKVTSGLGTVVDYKSLTKGLSIADIACAQYY 5086  
 NLSSV+P SR+AGRSA+ED+LFSK+VTSGLGTVD DYK+CTKGLSIADIACAQYY  
 Sbjct: 671 NLSSVIPSLPTSGSRVAGRSAIETDILFSKIVTSGLGTVDADYKNTCKGLSIADIACAQYY 730  
 70 Query: 5087 NGIMVLPGVADAERMAMYTGSLIGGMVLGGLTSAAAIPFSLAIQARLNVALQTDVLQEN 5266  
 NGIMVLPGVADAERMAMYTGSLIGG+ LGGLTSA +IPFSLA+QARLNVALQTDVLQEN  
 Sbjct: 731 NGIMVLPGVADAERMAMYTGSLIGGIALGGLTSAVSIPFSLAIQARLNVALQTDVLQEN 790  
 Query: 5267 QKILAASFNKAINNIVASFSSVNDAITHTAEIHTVTIALNKIQDVVNQQGSALNHLSQ 5446  
 QKILAASFNK+ NIV +F+ VNDAIT T++A+ TV ALNKIQDVVNQQG++LNHLSQ  
 Sbjct: 791 QKILAASFNKAMTNIVDAFTGVNDAITQTSQLQTVAALNKIQDVVNQQGNSLNHLSQ 850  
 75 Query: 5447 LRHNFOAISNSIHAIIYDRLDSIQADQQVDRLLITGRLAALNAFVSVQLNKYTEVRGSRRLA 5626

Sbjct: 851 LR NFQAISS+SI AIYDRLD+IQADQQVDRLITGRLAALN FVS L KYTEVR SR+LA  
 5 Query: 5627 QQKINECVKSQSNSRNGFCGNTHIFSIVNSAPDGLLFLHTVLLPTDYKNVKAWSGICVDG 5806  
 Sbjct: 911 QQKVNECVKSQSNSRNGFCGNTHIFSIVNAAPEGLVFLHTVLLPTQYKDVEAWSGLCVDG 970  
 10 Query: 5807 IYGYVLRQPNLVLYSDNGVFRVTSRVMFQPRLPVLSDFVQIYNCNVTVNISRVLHVTI 5986  
 Sbjct: 971 TNQYVLRQPNLALYKEGNYYRITSRIMFEPRIPTMADFVQIENCNVTVNISRSELQTTIV 1030  
 15 Query: 5987 PDYDVNKTLOEFAQNLPKYVKPNFDLTPFNLTYLNLSSELKQLEAKTA 6133  
 Sbjct: 1031 PEYIDVNKTLOELSYKLPNYTVPDLVVEQYNQTLNLSEISTLENKSA 1079

## 6. Sequence F

20 3062 Nucleotides encoding putative 3' end of Spike, hypothetical nsp 3, Envelope protein 5B, Matrix and  
 Nucleocapsid polypeptides  
 AGCTGATCGTTGATTGAGTTGCTTAATAGGTTTGAAGAAATTATATCAAATGGCCTGGTGGGTTGGCTCAT  
 TATTCTGTTGTTTGTGATTGAGCTCTCTGTGTTGTTGCTTCTACAGGTTGTTGGCTTGG  
 25 CAATTGTTAACCTCATCAATGCGAGGCTGTTGATTGTTCAACTAAACTTCTTATTATGAATTGAAAA  
 GGTCCACGTTCAATAATGCCTTCGGTGGCCTATTCAACTTACTCTGAAAGTACTATAAAGAGTGTGGCT  
 AATCTCAAATTACCACTCATGATGTTACTGCTCTGCGAACAATTAAACCTGTTACTACACTAGTACTATC  
 ACTGCTTATTGTTAGTTAGTTGCTCACTTATTGCTTATTCAAACCTCTTACTGCTAGAGGTCGTGTT  
 GCTTGTGTTGTTAAAACATTGACACTATCTGTCTATGTGCTTATTGGTTCTTGGTATGTATCTTGAC  
 AGTTTATAATTTTTCTACGCTGTTGATTCAATGTTGGCTATTATGCCATCTCTATAAAAATT  
 30 TTTCATTGTTTGTCAATGTTACTAAACTATGCTTCAGGCAAGTGTGTTGATCTGAACAATTCTT  
 ATGAAATCGTTTGTCTATTATGGTGGTGAACACTATGCTGTTTAGGTGGTGAACACTATTACTTTGTTT  
 CTTTGATGACCTTATGTTGCTATTAGAGGTCTTGTAAGAAACCTACAACCTATGCGTAAGGTTGACTTGT  
 ATAATGGTGTGTCATTACATTGGCCAGAGCCAGTTGTTGGTATAGTTTACTCCTCTCAACTACGAAG  
 ATGTTCTTCGATTAATTGATGACAATGGCATTGCTCTCAATTCTTATGGCTTGTGTTATGTT  
 35 TTTGTGTTGGCAATGACCTTATTAAACATTGCTATTGTTACTGCTTACTGCTATTGTTAGTAGGACATTA  
 TATCAACAGTTATAAAATTCTGCTTACCAAGATTATATGCAAATAGCACCTGCTCAGCTGAAGTACTA  
 AATGTCTAAACAGATGCTAAAGTGTGCTTACATTGTTAGGTTATGCTTACAGGTTTATGCTTACGTAACTGGAAC  
 TTAGTGGAAATTAAATTCTAACAGTTTATAGTGTGTTGAGTATGGCATTATAAGTATAGCAGACTCTTT  
 ATGGTTAAAGATGTCGTTTATGGTGTATGCCACTTGTCTAGCTTGTCTATTGACTGTTGTC  
 40 ATTAAATGTTGACTGGGCTTTGGTTTGTATTCTTATGCTTATTACACTTGTGTTATGGGTTATGT  
 ATTGTTAATAGTTGACTGGGCTTTGGGCTTAAACTTGGGCTTAAATCTGAAACTATGCAATCA  
 TCTCTCTCCAGGTTATGGACATAATTATTACCTACGGGCTGCTTGGCCTACAGGGTCACTTACACTTC  
 TTAGTGGTGTACTCTGTTGATGCCATTGGGCTTAAAGATTGCTACTGGTGTCAAGTGGGTCAGTGGCTTAAATATGAA  
 TAGTTGCTACACCTAGTACCAAACTGTTGACCGTGTGTTGGTGTGCTCTGTTAATGAAACACAAGCCAGACTGGTT  
 45 GGGCATTCTACGTCGCTAAACATGGTATTGCTGGTGTGCTCTCAGGAGGGTGTGTCAGAAAGAG  
 AGAAGTTGCTTCAATTAAACTAAACAAATGGTAGTGTAAATTGGCCGATGACAGAGCTGCTAGGAAG  
 AAATTCTCTCCCTCATTTACATGCCCTTTGGTTAGTGTGATAAGGCACCATATAGGGTCATTCCCGAG  
 AATCTGTCCTATTGGTAAGGGTAATAAAAGATGAGCAGATTGGTTATGGAATGTTCAAGAGCGTGGCGTATG  
 CGCAGGGGGCAACGTGTTGATTGCTCTAAAGTCAATTACCTAGGTACTGACCTCAAGGACCT  
 50 AAATTCAAGACAACGTTCTGATGGTGTGTTGGGTTGCTAAGGAAGGTGCTAAACTGTTAATACCAGTCTGGT  
 AATCGCAAACGTAATCAGAAACCTTGGGAAACCAAGTCTCTATTGCTTGGCTCCAGAGCTCTGTTGAG  
 TTTGAGGATGCTCTAAACTCATCTGCTAGCAGTGTCTTCAACTCGTAACAAACTCAGGAGACTCTTCT  
 CGTAGTACTTCAGACAACAGTCGCACTCGTCTGATTCTAACCAAGTCTCTCAGATCTGTTGCTGTT  
 ACTTGGCTTAAAGAACTTAGGTTGATAACCAGTCGAAGTCACCTAGTTCTGGTACTTCAACT  
 55 AAACCTAATAAGCCTCTTCTCAACCCAGGGCTGATAAGCCTTCTCAGTTGAAAGAAACCTGTTGGAAAGCGTGT  
 CCTACCAGAGAGGAAATGTTATTCACTGCTTGGTCTCGTGTATTAACTCACAATATGGGGATTCAGATCTT  
 GTTCAAGATGGTGTGATGCCAGGGTTTCCACAGCTTGTGATAATTGCTTACCTACAGGCTGCTTATTCTT  
 GATAGTGGGTTAGCAGTAAGTGGGTGATAATTGCTTACAGTACACTACAGGCTGCTTATTCTT  
 GATAATAAGAACCTCTTAAGTTCATTGAGCAGATTAGTGTCTTACTAAACCCAGTCTATCAAAGAAATGCA  
 60 TCAACATCATGTTGCTCAGAACACAGTACTTAATGCTTCTATTCCAGAACTAAACCATGGCTGATGAT  
 GATTCAAGCCATTATAGAAATTGTCAACGAGGTTTGCATTAAATTGTTGTAATTCCAGTTGAATGTTATTAT  
 TATTAGTTGCAACNCCCATGGTTAGGCCATGATAAGGGTTAGTCTACAAACGATCAAGCT

### 65 Hypothesised ORFs

>-out: 17 to 238: Frame 2 74 aa  
 FELLNRFENYIKWPWWVWLIIISVVFVLLSLLVFCCLSTGCCGCNCLSSMRGCCDCGSTKLPPYYEFKVVHQ  
 >-out: 223 to 723: Frame 1 167 aa

KGPRSIMPFGGLFQLTLESTINKSVANLKLPPHDVTVLRDNLKPVTTLSTITAYLLVSLFVTVYFALFKPLTAR  
 VACFVULKLLTLSVYVPLLVFGMYLDSFIIFFLRCCFDSYMLAIMPIKIFHLFCMMLNYASFQASVGLNT  
 FMKIVLILLFMVVITMSF  
 >-out: 525 to 917: Frame 3 131 aa  
 5 QFYNNFFSTLLFRFIHVGYYAYLYKNFSFVLFNFVTKLCFVSGKCWYLEQSFYENRFAAIYGGDHVVVLGGETITI  
 SFDDLYVAIRGSCEKNLQLMRKVVDLYNGAVIYIFAEPPVVGIVYSSQLYEDVPSIN  
 >-out: 877 to 1131: Frame 1 85 aa  
 10 FTPLNYTKMFLRLIDDDNGIVLNSILWLLVMIFFVLAATFIKLIQLCFTCHYFFSRTLYQPVYKIFLAYQDYM  
 APVPAEVLNV  
 >-out: 1140 to 1820: Frame 3 227 aa  
 15 TMSNSSVPLSEVVVHRLRNWNFSWNLILTVFIVVLYQYGHYKYSRLLYGLKMSVWLWPLVLALSIFDCFVNFN  
 WVFFGFSILMSIITLCLWVMYFVNSFRLWRRVKTFWAFNPETNAIIISLQVYGHNYYLPMVAAPTGVTLTLLSGV  
 LVDGHKIA TRVQVQQLPKYVIVATPSTIVCDRVGRSVNETSQTGWAFYVRAKGDFSGVASQEGVLSEREKLI  
 LI  
 >-out: 1324 to 1539: Frame 1 72 aa  
 20 LCLFLTVLSILMWTSFVLFVLCLLLHFVYGLCILLIVSDFGAVLKLFGLLILKLMQSSLSRFMDIIITYR  
 >-out: 1654 to 1815: Frame 1 54 aa  
 LLHLPQLFVTVLVALLMKQARLGHSTSVLNMVIFLVLPLRRVFCQKERSCFI  
 >-out: 1819 to 2964: Frame 1 382 aa  
 25 SKLNKMASVNWADDRAARKKFPPPSFYMPPLLSSDKAPYRVI PRNLVPIGKGNKDEQIGYWNVQERWRMRRGQR  
 DLPPKVHFYILGTGPHKDLKFRQRSDGVVVAKEGAKTVNTSLGNRKRNQKPLEPKFSIALPPELSVVEFEDRS  
 NSSRASSRSSTRNNSRDSSRSTSRRQSRTRSDSNQSSSDLVAAVTLALKNLGFNDNQSKSPSSGTSTPKPNKE  
 SQPRADKPSQLKKPRWKRVPTREENVICQCFGPRDFNHNMGDSLVLQNGVDAKGFQQLAELIPNQAALFFDSEVS  
 DEVGDNVQITYTYKMLVAKDNKNLPKFEQISAFTPSSIKEMQSQS SHVAQNTVLANASIPESKPLADDDSAII  
 IVNEVLH  
 >-out: 1847 to 2074: Frame 2 76 aa  
 30 IGPMTTEILLGRNFLLHFTCLFWLVLIRHHIGSFPGILSLLVRVIKMSRLVIGMFKSVGVCAGGNVLICLLKFIF  
 T  
 >-out: 2078 to 2410: Frame 2 111 aa  
 VLDLIRTLNSDNVLMVLFGLLRKVLKLLIPVLVIANVIRNLWNQSSLLLCLQSSLLLRLIALITHLVLAVVLQ  
 VTTHETLLVVLQDNSLALVLLTSLQI LLLLWL  
 >-out: 2771 to 2938: Frame 2 56 aa  
 35 LRIIIRTFLSSLSRLVLLNPVLSKKCSHNHMLRTQYLMLLFQNLNHWLMMI QPL

### Alignment

>gi|13604396|gb|AAK32190.1| spike glycoprotein [Human coronavirus 229E]  
 40 Length = 1173  
 Score = 50.4 bits (119), Expect = 7e-06  
 Identities = 26/71 (36%), Positives = 31/71 (43%)  
 Frame = +2

45 Query: 26 LNRFENYIKWPWXXXSMRGCCDCGSTKL 205  
 LNR E YIKWPW S+RGCC+ STKL  
 Sbjct: 1105 LNRVETYIKWPWWVWL CISVVLIFVVSMLLCCCSTGCCGFSCFASSIRGCCE--STKL 1162

50 Query: 206 PYYEFEKVHVQ 238  
 PYY+ EK+H+Q  
 Sbjct: 1163 PYDVEKIHIQ 1173

55 >gi|12175749|ref|NP\_073552.1| 4a protein [Human coronavirus 229E]  
 gi|138983|sp|P19739|VN4A CVH22 Nonstructural protein 4a (ORF4a)  
 gi|74871|pir|1MNIHHC nonstructural protein 4 - human coronavirus (strain 229E)  
 gi|58928|emb|CAA33682.1| unnamed protein product [Human coronavirus 229E]  
 gi|12082742|gb|AAG48598.1| 4a protein [Human coronavirus 229E]  
 Length = 133

60 Score = 71.6 bits (174), Expect(2) = 1e-17  
 Identities = 41/95 (43%), Positives = 56/95 (58%)  
 Frame = +1

65 Query: 253 GLFQLTLESTINKSVANLKLPPHDVTVLRDNLKPVTTLSTITAYLLVSLFVTVYFALFKPL 432  
 GLF I L S +N+S++N K+ + ++K T + AY L+SLFV YFALFK  
 Sbjct: 4 GLFTIQLVSAVNQSLSNAKVSAEVSRQVIQDVKDGTVTFNLLAYTLMMSLFVVYFALFKAR 63

70 Query: 433 TARGRVACFVLKLLTLSVYVPLLVLFGMYLDSFII 537  
 + RGR A V K+L L VYVPLL Y+ + +I

## Fig 3. (Cont.)

24/25

Sbjct: 64 SHRGRAALIVFKILILFVYVPLLYWSQAYIYATLI 98

5 Score = 40.4 bits (93), Expect(2) = 1e-17  
 Identities = 15/30 (50%), Positives = 22/30 (73%)  
 Frame = +3

10 Query: 549 LLFRFIHVGGYYAYLYKNFSFVLFNVTKLCF 638  
 LL RF H ++ +LYK + F++FNVT LC+  
 Sbjct: 102 LLGRFFHTAWHCWLKTYKTWDFIVFNVTLCY 131

15 >gi|12175750|ref|NP\_073553.1| 4b protein [Human coronavirus 229E]  
 gi|188992|sp|P19740|VN4B\_CVH22 Nonstructural protein 4b (Nonstructural protein 5A) (ORF4b)  
 gi|74872|pir|MNIIHH2 nonstructural protein 5A - human coronavirus (strain 229E)  
 gi|58924|emb|CAA33683.1| unnamed protein product [Human coronavirus 229E]  
 gi|12082743|gb|AAG48594.1| 4b protein [Human coronavirus 229E]  
 20 Length = 88  
 Score = 86.7 bits (213), Expect = 2e-16  
 Identities = 38/80 (47%), Positives = 54/80 (67%)  
 Frame = +1

25 Query: 640 VSGKCWYLEQSFYENRFAAIYGGDHYVVLGGETITFVFSFDDLYVAIRGSCEKNLQLMRKV 819  
 + GKCW+LE + F YGGD ++ +G +++ S +DLYVA+RG +K+L L RKV  
 Sbjct: 1 MQGKCFWLENKALKP-FVCFYGGDQFLYIGDRIVSYFSTNDLYVALRGRIDKDLSSLRKV 59

30 Query: 820 DLYNGAVIYIYFAAEPVVGIV 879  
 +LYNG +Y+F E P VGIV  
 Sbjct: 60 ELYNGECVYLFCCEHPAVGIV 79  
 >gi|12175751|ref|NP\_073554.1| envelope protein [Human coronavirus 229E]  
 35 gi|188994|sp|P19741|VEMP\_CVH22 Envelope protein (Protein 5B)  
 gi|74873|pir|MNIIHH3 nonstructural protein 5B - human coronavirus (strain 229E)  
 gi|58925|emb|CAA33684.1| unnamed protein product [Human coronavirus 229E]  
 gi|12082744|gb|AAG48595.1| envelope protein [Human coronavirus 229E]  
 40 Length = 77  
 Score = 87.8 bits (216), Expect = 3e-17  
 Identities = 36/76 (47%), Positives = 55/76 (72%)  
 Frame = +3

45 Query: 901 MFLRLIDDDNGIVLNSILWLLVMIFFFVILAMTFIKLIQLCFTCHYFFSRTLYQPVYKIFLA 1080  
 MFL+L+DD+ +V+N +LW +V+I ++ +T IKLI+LCFTCH F +RT+Y P+ ++  
 Sbjct: 1 MFLKLVDDHALVVNVLLWCVVLIVILLVCITIILKLIKLCFTCHMFCNRTVYGPPIKNVYHI 60

50 Query: 1081 YQDYMQIAPVPAEVLN  
 YQ YM I P P V++  
 Sbjct: 61 YQSYMHIDPFPKRVID 76  
 >gi|74837|pir|MMIHHC E1 membrane glycoprotein - human coronavirus (strain 229E)  
 55 gi|329573|gb|AAA45461.1| membrane protein [Human coronavirus 229E]  
 Length = 225  
 Score = 275 bits (703), Expect = 4e-72  
 Identities = 128/224 (57%), Positives = 159/224 (70%)  
 Frame = +3

60 Query: 1143 MSNSSVPLSEVYVHLRNWNFSWNLILTVFIVVLQYGHYKYSRLLYGLKMSVLWCLWPLVL 1322  
 MSN + ++ HL+NWNF WN+ILT+FIV+LQ+GHYKYSRLLYGLKM VLW LWPLVL  
 Sbjct: 1 MSNDNCT-GDIVTHLKNWNFGWNVILTIIFIVILQFGHYKYSRLLYGLKMLVLWLLWPLVL 59

65 Query: 1323 ALSIFDCFVNFDWVWVPGFSIILMSIITLCLWVMYFVNSFRLWRRVKTWFAFNPETNAAI 1502  
 ALSIFD + N++ +W F FS+LM++ TL +WVVF NSFRL+RR +TFWA+NPE NAI  
 Sbjct: 60 ALSIFDTWANWDSNWAFAVAFSLLMAVSTLVMWVMYFANSFRLFRRARTFWAWNPEVNAIT 119

70 Query: 1503 SLQVYGHNYYLPMMAAPXXXXXXXXXXXXXXHKIATRVQVQLPKYVIVATPSTTIIVC 1682  
 V G YY P+ AP H++A+ VQV LP+Y+ VA PSTTI+  
 Sbjct: 120 VTTVLGQTYYQPIQQAPTGITVLLSGVLYVDGHLASGVQVHNLPEYMTVAVPSTTIY 179

Query: 1683 DRVGRSVNETSQTGWFYVRAKHGDFSGVASQEGVLSEREKLLH 1814  
 RVGRSVN + TGW FYVR KHGDFS V+S ++E E+LLH  
 Sbjct: 180 SRVGRSVNSQNSTGWVFYVRVKHGDFAVSSPMNSMTENERLLH 223

5 >gi|12175758|ref|NP\_078556.1| nucleocapsid protein [Human coronavirus 229E]  
 gi|29840828|sp|P15190|NCAP\_CVH22| Nucleocapsid protein (N structural protein) (NC)  
 gi|77063|pir||S08031| nucleocapsid protein - human coronavirus  
 gi|58933|emb|CAA35708.1| unnamed protein product [Human coronavirus 229E]  
 gi|12082746|gb|AAG48597.1| nucleocapsid protein [Human coronavirus 229E]

10 Length = 389

Score = 267 bits (682), Expect = 1e-69  
 Identities = 159/406 (39%), Positives = 222/406 (54%), Gaps = 31/406 (7%)  
 Frame = +1

15 Query: 1834 MASVNWAD---DRAARKKFPPPSFYMPPLLVSSDKAPYRVIPRNLVPIGKGNKDEQIGYWN 2004  
 MA+V WAD + R+ P S Y PLLV S++ P++VIPRNLVPI K +K++ IGYWN  
 Sbjct: 1 MATVKWADASEPQRGRQGRIPYSLYSPLLVDSEQ-PWKVIPRNLVPINKDKNKLIGYWN 59

20 Query: 2005 VQERWRMRRGQRVDLPPKVHFYYLGTGPHKDLKFRQRSDGVVVVAKEGAKTVNTSLGNRK 2184  
 VQ+R+R R+G+RVDL PK+HFYYLGTGPHKD KFR+R +GVVVVA +GAKT T G R+  
 Sbjct: 60 VQKRFTRKGKRVDLSPKLHFYYLGTGPHKDAKFRERVEGVVVAVDGAKTEPTGYGVRR 119

25 Query: 2185 RNQKPLEPKFSIALPPELSVVEFEDXXXXXXXXXXXXXXXXXXXXXXXXXXXX 2364  
 +N +P P F+ LP ++VVE D  
 Sbjct: 120 KNSEPEIPHFNQKLPNGVTVVEPD----SRAPSRSQSRSQSRSRGRGESKPQSRNPSSDR 174

Query: 2365 XXXXXXXLVAAVTLALKNLGFDN-----QXXXXXXXXXXXXXXXXXXXX 2496  
 ++ AV ALK+LGFD  
 Sbjct: 175 NHNSQDDIMKAVAAALKSLGFDKPKQEKDKSAKTGTPKPSRNQSPASSQTSAKSLARSQS 234

Query: 2497 QPRADKPSQLKKPRWKRVPTRE--ENVIQCFGPRDFNHNMGDSDLVQNGVDAKGFPQLAE 2670  
 ++ +++KPRWKR P + NV QCFGPRD +HN G + +V NGV AKG+PQ AE  
 Sbjct: 235 SETKEOKHEMQKPRWKRQPNDDVTSNVTQCFGPRDLDHNFGSAGVVANGVAKGYPQFAE 294

35 Query: 2671 LIPNQAALFFDSEVSTDEVGDNVQITYTYKMLVAKDNKNLPKFIEQISAFTKPSSIKEMQ 2850  
 L+P+ AA+ FDS + + E G+ V +T+T ++ V KD+ +L KF+E+++AFT +EMQ  
 Sbjct: 295 LVPSTAAMLFDSHIVSKESGNTVVLTFTRVTVPKDHPHLGKFLEELNAFT----REMQ 349

40 Query: 2851 SQSSHVAQNTVLNASIPE-----SKPLADDDSAIIIEIVNEV 2958  
 Q+ +LN S E ++P+ D+ S +I++EV  
 Sbjct: 350 -----QHPLLNP SALEFNPSQTSPATAEPVRDEVSIETDIIDEV 388

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